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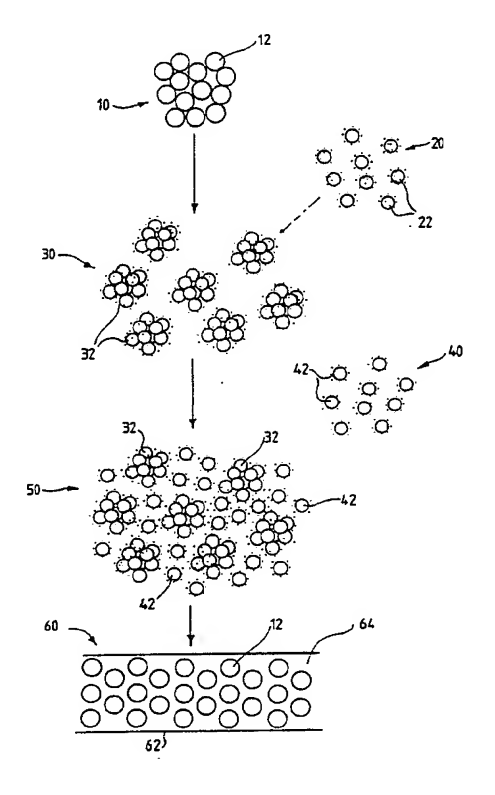
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(54) Title: AQUEOUS DISPERSION OF COMPOSITE PARTICLES INCLUDING POLYMERIC LATEX

(57) Abstract

Inorganic material particles such as titanium dioxide (10) pigment particles (12) are dispersed in an aqueous medium with a polyelectrolyte pigment dispersant (20). A selected, relatively hard, polymeric latex (40), which adsorbs onto the surface of the inorganic material particles in the presence of the pigment dispersant, is added, as well as a film-forming binder (64) to give a formulated coating composition. The strongly adsorbing polymeric latex can include residues of polymerized itaconic acid or dihydrogen phosphate functional groups, or have a high level of acid functional groups or acid functional groups distributed preferentially at the surface of the latex. The adsorbed latex particles space the inorganic particles apart in the coating film (60), improving performance properties. Alternatively, a single selected polymeric latex is used to both strongly adsorb to the pigment particles and to serve as binder, or a mixture of a selected, strongly adsorbing polymeric latex having a low effective glass transition temperature and a conventional binder latex is employed.







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AQUEOUS DISPERTION OF COMPOSITE PARTICLES INCLUDING POLYMERIC LATEX.

BACKGROUND OF THE INVENTION

1. Field Of The Invention

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This invention relates generally to a process for preparing aqueous compositions including inorganic material particles, such as aqueous compositions including titanium dioxide pigment, and more particularly to the distribution of inorganic material particles in products formed by the loss of water from such aqueous compositions.

2. Background Of The Invention

Aqueous dispersions of polymeric latex are used to 10 prepare a tremendous variety of commercially important products, including paints, coatings, primers, mastics, caulks, binders for non-woven materials, and adhesives. Frequently, the polymeric latex is included to form a continuous phase as the water is lost in order to bind the 15 product together, and to contribute important physical properties. In addition to the polymeric latex, most products include particles of one or more inorganic materials. Some inorganic materials contribute an important functional property to the product, such as fire resistance, 20 sheen, flatting, hardness, color, texture, opacity, or toughness. Often these inorganic materials are relatively expensive, and it is desirable to employ them in the formulated product as efficiently as possible. In other instances, the inorganic materials, for example, fillers, 25 may be less expensive than the latex and it may be desired to incorporate the maximum amounts possible without compromising the desirable properties attributable to the latex, such as, for example, binding and tensile properties,

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such as elongation and, in the case of adhesives, minimum loss of tackiness. Since the polymeric latex is typically the most expensive component of the formulated product, inexpensive inorganic materials are frequently included to reduce product cost. In this case, it is desirable to use the greatest possible volume of the inexpensive inorganic material, while still retaining the desired performance properties of the product, since these properties typically decline when too much of the inexpensive inorganic material is included. A good example is a paint formulated with so much of an inexpensive filler pigment, such as calcium carbonate, that the critical pigment volume concentration is exceeded. Such a paint will provide a much less protective coating for a surface on which it is applied than a similar paint formulated with slightly less filler, and having a pigment volume concentration below the critical level.

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One way of making efficient use of the inorganic material employed is by dispersing this pigment as well as possible. Agglomerated or aggregated inorganic material pigment particles frequently make less than their full potential contribution to the performance of the product, such as, for example, strength and resistance properties. Inorganic materials are typically sold as dry powders. As a practical matter, this powder must be milled in a liquid medium to break up agglomerates and to obtain a colloidally stable dispersion. Milling conventionally takes place in the presence of a polyelectrolyte particle dispersant which adsorbs to the particle surface.

Titanium dioxide has been for many years the pigment of choice for conferring opacity to plastic sheets and films, and particularly to coatings formed from coating compositions and paints. In paints titanium dioxide is typically the most expensive component of the formulation. It has been an ongoing objective of the coatings industry to achieve the desired degree of opacity in a coating while at the same time using as little titanium dioxide pigment as

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possible. One way in which this is done is by employing titanium dioxide which has an optimal average particle size and particle size distribution for scattering light.

Another way of making efficient use of the titanium dioxide employed is by dispersing this pigment as well as possible.

Agglomerated or aggregated titanium dioxide pigment particles make less than their full potential contribution to the performance of the coatings, such as, for example, with regard to coating opacity and colorant potential. Such aggregates can also impair certain other properties of the coating or film such as, for example, strength and resistance properties. Titanium dioxide is often sold as a dry powder. As a practical matter, this powder must be milled in a liquid medium to break up agglomerates and to obtain a colloidally stable dispersion.

In order to stabilize the product formulation against settling or flocculation, a variety of surface active dispersing additives have been used. Manufacturers have often used inorganic materials as dry powders. In this case, a dispersing additive is usually added directly to the material and a small amount of water in a preliminary "grind" step in which loose agglomerates of the inorganic material particles are broken up by mechanical shear forces. The dispersing additive typically remains in the mix through the subsequent steps in the product preparation process and typically will be found in the fully formulated product composition.

Sometimes inorganic materials are commercially

supplied in concentrated slurry form, such as, for example, in water. However, since the inorganic material particles in the slurries are prone to aggregation upon storage, the slurries often need to be redispersed for maximum effectiveness before use in formulating a coating

composition. Either the manufacturer of the inorganic material slurry or the end user of the slurry, and sometimes

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both, may perform the redispersion step.

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The opacifying capability or hiding power of a paint or coating is a function of the spacing of the titanium pigment particles in the dried coating. The light scattering characteristics of titanium dioxide particles are well known. The average size and size distribution of titanium dioxide particles used for opacifying purposes has been highly optimized by the titanium dioxide manufacturers for maximum scattering. Maximum light scattering occurs when the titanium dioxide pigment particles have a diameter of from about 200 to about 250 nanometers and are spaced far apart from each other, on the order of a few particle diameters, so that there is minimal interference between the light scattering of neighboring particles.

In practice, however, for example in the formulation of paints, it is widely recognized that when enough titanium dioxide particles have been dispersed into the polymeric coating vehicle to yield films of acceptable opacity, the level of opacity which is achieved is significantly less than would be theoretically predicted from the light scattering potential of a single titanium dioxide particle multiplied by the total number of particles in the path of light passing through the film.

A number of factors have been identified which partially account for the diminution of opacity from the theoretical predictions. If two or more titanium dioxide particles are in actual contact with each other, or even if they are closer together than the optimum spacing distance, the particles will behave like a single aggregated larger particle and result in reduced light scattering. This occurs if the titanium dioxide particles are not adequately deaggregated during the dispersion process. However, even if the titanium dioxide particles are fully deaggregated in the dispersion process, a random distribution of particles will not provide the maximum scattering achievable in theory if the particles were optimally distributed.

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In a related phenomenon, referred to as "crowding," titanium dioxide pigment particles are forced to be nearer to each other than the desired optimum merely by the lack of available space. This lack of available space may be caused by the space taken up by the other coating fillers and extenders which are of a comparable size to, or which are larger than, the pigment particles. In the case of polymeric binders, which are particulate in nature prior to film formation, such as, for example, latex or emulsion polymers, and in the case of nonaqueous dispersion polymers, 10 the binder polymer particles themselves can crowd pigment particles, especially if they are of comparable or larger in size than the titanium dioxide particles.

The traditional guiding rule or goal in the formulation of practical, dispersed titanium 15 dioxide-containing coatings, such as, for example, pigmented latex paints, is to make the titanium dioxide dispersion and the polymeric latex binder dispersions as colloidally stable and compatible with each other as possible. Nevertheless, in coatings formed from such dispersions the distribution of 20 the titanium dioxide particles in the polymer system at best approaches that of a random distribution. As a result, there exist a substantial number of titanium dioxide particles in close proximity to each other, and possibly in direct physical contact with each other. 25

In the case of latex paints and coatings, it is conventional practice to first form a stable aqueous dispersion of titanium dioxide pigment with other fillers or extenders. This dispersion, also known as a "mill base" or "grind," may contain water-miscible solvents, such as for example glycols and glycol ethers, and relatively low molecular weight water soluble polyelectrolytes as titanium dioxide pigment grinding aids or dispersants. Generally, these pigment dispersants are anionic polyelectrolytes. Many different types of such dispersants are known. For

35 example, U.S. Patent 2,930,775 discloses the water soluble

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salts of diisobutylene naleic anhydride copolymers having molecular weights between about 750 and 5,000 as dispersants when employed at concentrations of from about 0.05 to 4% on pigment weight. U.S. Patent 4,102,843 and U.S. Reissue Patent 31,936 disclose the use of water soluble salts of copolymers of hydroxyalkyl-(meth)acrylates and (meth)acrylic acid of molecular weights of from 500 to 15,000 at concentrations of from about 0.01 to 5% on pigment to produce glossy emulsion paints. U.S. Patent 4,243,430 discloses a water-soluble dispersant comprising an addition 10 copolymer comprising greater than 30% alpha, betaunsaturated monovinylidene carboxylic acid, the copolymer having an apparent pKa between 6.0 and 7.5 and molecular weight between about 500 and 15,000, and forming a water soluble salt with zinc ammonia complex ion. Low molecular 15 weight polyphosphate salts, such as potassium tripolyphosphate, are also used because they are relatively inexpensive, but they tend to have marginal hydrolytic stability.

The use of these and other polyelectrolyte dispersants is described in T.C. Patton, Paint Flow and Pigment Dispersion, (Wiley Interscience, 2nd edition) 290-295 (1979). Also described therein (pages 468-497) are a number of milling devices used in the preparation of pigment dispersions. One such device which is commonly used in the 25 manufacture of latex paints is the highspeed disk disperser designed to develop high shearing forces in the pigment grinding step. Common practice is to use the device with dispersant to form a stable dispersion of titanium dioxide pigment, and then to add to the dispersion the aqueous latex 30 polymer binder along with the other ingredients, such as for example thickeners and rheology modifiers, antifoaming agents, colorants, coalescing agents or temporary plasticizers for the latex polymer particles, and surfactants for substrate wetting and colorant 35 compatibility. The pigment dispersion process and the

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relation to flocculation and optical properties are discussed further in <u>Treatise on Coatings</u>, Vol. III, Part 1 (Marcel Decker) (1975); <u>Pigment Handbook</u>, Vol. I (2nd edition, Wiley Interscience) (1988), and <u>Pigment Handbook</u>, Vol. III (Wiley Interscience) (1973).

There is a continuing need to improve the effective utilization of inorganic materials such as titanium dioxide in aqueous coating compositions and thereby to improve the opacity and other performance properties of coating compositions.

SUMMARY OF THE INVENTION

The present invention provides a process for preparing an aqueous dispersion of composite particles, the composite particles each including a plurality of polymeric latex particles adsorbed onto an inorganic material particle, such as a titanium dioxide particle. The resulting composite particles provided by the present invention can be used in preparing formulated aqueous compositions, such as coating compositions, adhesives, binders, for non-woven materials, and paints, which in turn give coatings and other products in which the inorganic material is more efficaciously dispersed and distributed than in prior art formulations.

The process of the present invention contemplates dispersing inorganic material particles such as titanium dioxide pigment particles in an aqueous medium with a pigment dispersant, such as a polyelectrolyte pigment dispersant. The process of this invention further contemplates preparing a dispersion or suspension of selected polymeric latex particles in the aqueous medium. These polymeric latex particles are selected to adsorb onto the dispersed inorganic pigment particles to provide composite particles, each including both an inorganic material particle and adsorbed polymeric latex particles. The inorganic material particles remain dispersed in the

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aqueous medium during and after adsorption of the polymeric latex particles, that is, during their transformation into composite particles. The selected polymeric latex particles are believed to strongly adsorb onto the inorganic material particles. In some cases, the selected polymeric latex particles are believed to, in effect, irreversibly adsorb onto the inorganic material particles. In at least some cases, it is believed that the previously adsorbed pigment dispersant is displaced by the adsorbed polymeric latex particles.

The selected polymeric latex particles can be prepared by processes which are believed to provide a high concentration of acid residues at the surface of the particles, processes which employ a high level of acid-functional monomer, or processes which provide copolymerization of adsorption-promoting monomers, such as dihydrogen phosphate-functional monomers or dicarboxylic acid monomers such as itaconic acid.

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In addition to the selected polymeric latex, other 20 components can be added depending on the application for the aqueous composition. For example, a film-forming binder such as a relatively soft polymeric latex can be included, as well as other onventional components used to provide a formulated coating composition, such as coalescents, preservatives, thickeners, and rheology control agents. The adsorbed latex particles remain adsorbed during the film-formation, loss of water and other fugative components, and serve to space the inorganic particles apart in the coating film, improving performance properties.

Preferably, the selected polymeric latex particles are polymerized from monomer which provides polymer which is hard or rigid at the temperature at which the composite particles are to be used, such as monomer which provides a polymeric material with an effective glass transition temperature of at least about 20 °C, more preferably at least about 35 °C, and even more preferably at least about

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50 °C in the case of a composition including composite particles and applied at ambient or room temperature (that is, at about 20 - 23 °C). Higher temperature ranges may be more appropriate for applications at higher temperatures,

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baked coatings. The rigidity of the adsorbed polymeric latex particles is believed to aid in spacing adjacent inorganic material particles.

Alternatively, the selected polymeric latex can have a low effective glass transition temperature and serve as a binder for the composition. However, softer polymeric latex particles are thought to deform during film-formation, such that somewhat larger particles may be required to obtain the same pigment particle spacing as with a hard or rigid polymeric latex.

The improved dispersion of the inorganic material particles provided by the present invention advantageously provides greater flexibility in preparing aqueous compositions including dispersed inorganic material. For example, in some cases the present process provides a means of reducing the amount of inorganic material required to provide desired properties, such as mechanical properties or opacification. When the composite particles include titanium dioxide and are used in coating compositions, the present process provides improved coating properties, such as opacity or hiding, tint strength, mechanical properties, "ICI" or "cone-and-plate" (high shear rate) viscosity, gloss, and scrub resistance. The process also provides coating compositions giving coatings with improved opacity for a given pigment level.

The composite particles resulting from the process of this invention are particularly useful for improving the performance properties and permeability of coatings and paints formed therefrom. Alternatively, the process of the invention offers the ability to formulate coatings and films of substantially equal performance properties as

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conventional systems but with lower concentrations of expensive inorganic material particle concentrations, such as in the case of titanium dioxide, and accordingly at lower cost. Similarly, higher concentrations of low cost inorganic materials can be obtained using the process of the present invention.

The pigment dispersant can be a polyelectrolyte dispersant. These polyelectrolyte dispersants, such as potassium tripolyphosphate, polyacrylic acid, polymethacrylic acid, and the like, which are used as 10 dispersion aids in preparing a pigment grind for a coating composition, are believed to adsorb to the surface of the inorganic materials in the grind, including any titanium dioxide particles, and thus increase the absolute value of the surface charge density or surface potential of the 15 pigment particles. As polymeric latex particles are frequently stabilized in aqueous media by surface charge of the same sign (negative) as the polyelectrolyte dispersants typically exhibit, the effect of employing polyelectrolyte 20 dispersants can be to increase electrical repulsive forces between inorganic material particles and polymeric latex particles. Adsorption of polymeric latex particles onto the inorganic material particles would appear less likely when the inorganic material particle is stabilized by a polyelectrolyte dispersant. Thus, the extent of adsorption 25 of the selected polymeric latex particles of the present invention is unexpected.

The process of the present invention further contemplates preparing fully formulated aqueous compositions, including aqueous coating compositions, using the composite particles formed by adsorption of the polymeric latex particles onto the inorganic material particles, and the subsequent use of the fully formulated aqueous compositions to form products, including coatings, and coated articles. While the aqueous medium containing the composite particles can be used directly in

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some cases to form products, in many instances it is desirable to employ the aqueous medium including the composite particles as an intermediate in the production of an aqueous composition, such as a coating composition, including one or more additional components (a "fully formulated" composition).

In one presently preferred embodiment, the process of this invention contemplates preparing a mixture of at least two types of polymeric latex particles, the first type being the selected polymeric latex particles to adsorb to the dispersed inorganic material particles, and preferably having an effective glass transition temperature of at least about 20 °C, more preferably at least about 35 °C, and even more preferably at least about 50 °C. The second type of polymeric latex particle is provided toform a binder for the composite particles. The mixture can be used to prepare formulated compositions, such as coating compositions, in which inorganic particles are advantageously spaced by adsorbed selected polymeric latex particles to provide improved performance properties.

In another preferred embodiment, two or more types of soft polymeric particles are employed as binders, one or more of these being a selected polymer latex which strongly adsorbs to the inorganic material particles.

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BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 is a schematic illustration of a preferred embodiment of the process of the present invention.

Fig. 2 is a schematic illustration of the distribution of titanium dioxide particles in a coating film formed from an aqueous composition prepared using a conventional prior art process.

Fig. 3 is a schematic illustration of the distribution of titanium dioxide particles in a coating film formed from an aqueous coating composition prepared using the process of

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the present invention.

Fig. 4a is a scanning electron micrograph showing composite particles prepared by the process of this invention.

Fig. 4b is a scanning electron micrograph showing a mixture of polymeric latex particles and inorganic material particles prepared by a prior art process.

DETAILED DESCRIPTION

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The process of the present invention provides selected polymeric latexes for use in adsorption onto inorganic material particles, the inorganic material particles having been initially dispersed using a pigment dispersant, such as a polyelectrolyte dispersant, or a steric stabilization agent.

The process provides a solution to the problem of how to practically utilize polymeric latex particles and inorganic material particles to improve the distribution and stability of the inorganic material inorganic material particles in aqueous-based coatings containing high concentrations of inorganic material particles.

Polymeric latex particles of selected composition, size and surface charge can be effectively used in the process of the present invention along with inorganic material particles, including in particular titanium dioxide particles, in concentrated dispersions. The selected polymeric latex particles function in the process by adsorbing onto the surface of the inorganic material particles in the presence of the initially adsorbed pigment dispersant. While not being bound by any particular theory of the molecular mechanism of the process of the present invention, it is presently believed that the initial pigment dispersant adsorbed to the surface of the inorganic material particles is in equilibrium with initial pigment dispersant in the aqueous medium, that the initial pigment dispersant

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does not completely cover the surface of the inorganic material particles at all times, and that at least a portion of the surface of the inorganic material particle is available for adsorption of the selected polymeric latex particles.

The preferred polymeric latex particles are believed to adsorb essentially irreversibly on the surface of the inorganic material particles, so that in time the initial pigment dispersant is replaced at least to some extent on the surface by the polymeric latex. The polymeric latex is preferably selected to provide this strong adsorption. This can be accomplished in different ways. In one presently preferred embodiment, a polymeric latex is polymerized from monomer mixture including a high level such as at least about ten percent by weight of polymer solids of carboxylic acid-functional monomer. In another presently preferred embodiment, a polymeric latex having acidic functional groups preferrentially distributed proximate the surface is employed. In still another presently preferred embodiment, a polymeric latex prepared from monomer including at least one polymerizable ethylenically unsaturated dihydrogen phosphate ester, such as a dihydrogen phosphate ester of 2-hydroxyethyl methacrylate, is employed. In yet another presently preferred embodiment, the polymeric latex is polymerized from monomer including a dicarboxylic acid monomer, such as itaconic acid.

As distinguished from prior art processes in which polymerization takes place in the presence of and/or on the surface of inorganic particles, the process of present invention utilizes selected, separately formed, polymeric latex particles to improve the distribution of concentrated inorganic material particles in polymer compositions, including dried films formed from aqueous coating compositions. More particularly, the process provides means for improving the distribution of inorganic material particles, and especially titanium dioxide pigment articles.

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such as those present at concentrations of at least about three percent by volume of the nonvolatile components of an aqueous composition, by adsorption of at least one selected, separately polymerized, polymeric latex onto the inorganic material particles, thereby forming a stable dispersion of microcomposite particles.

The present process provides for the adsorption of the selected polymeric latex on the surface of the inorganic material particles in the presence of the initial pigment dispersant, in concentrated, stable dispersions. The composite particles resulting from this adsorption process are particularly useful for improving the opacity, gloss, high shear viscosity, color development, permeability, and mechanical properties of coatings and films formed therefrom. Alternatively, the adsorption process of the invention offers the ability to formulate coatings and films of substantially equal performance properties as conventional systems but with lower inorganic material particle concentrations and accordingly at lower cost. In 20 addition, the present invention provides the opportunity to formulate compositions using greater quanities of low cost inorganic material particles than otherwise, while yet providing good performance properties.

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The selected polymeric latex particles of the present invention adsorb onto the surface of the dispersed inorganic material particles. Preferably, the selected polymeric latex particles adsorb strongly. There are many prior art processes in which dispersed inorganic material particles and polymeric latex particles are present together in an aqueous medium. For example, many aqueous coating compositions and water-based paints include such inorganic material particles and polymeric latex particles, the polymeric latex particles serving to form the binder for the inorganic material particles in a coating formed from the composition. However, in these compositions, the polymeric latex particles do not adsorb strongly to the dispersed

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inorganic material particles, as both the inorganic material particles and the polymeric latex particles will typically have surface potentials with the same sign, typically negative, the like electrical charges serving to stabilize the system by mutual repulsion.

One embodiment of the process of the present invention is illustrated schematically in Figure 1. A powder of titanium dioxide pigment 10 including titanium dioxide particles 12, partially aggregated or agglomerated (by van der Waals, or contact forces, or by moisture, or the like), 10 is mixed with a solution of a polyelectrolyte dispersant 20. Typically, an amount of an aqueous medium (not shown) is added to reduce the concentration of the particles. The mixture of titanium dioxide pigment 10 and the polyelectrolyte dispersant 20 is subjected to shear forces 15 in a disperser (not shown), and the polyelectrolyte dispersant macromolecules 20 spontaneously adsorb to the surface of titanium dioxide particles 12 to form a slurry or pigment grind 30 of dispersed titanium dioxide particles 32 in the aqueous medium. The adsorbed polyelectrolyte 20 dispersant increases the absolute value of the surface potential of the titanium dioxide particles. Without the addition of the polyelectrolyte dispersant and the application of shear forces by mixing, the titanium dioxide particles may form homo-aggregates, as conditions are 25 preferably selected so that the titanium dioxide particles are un-stable with respect to homo-coagulation.

Figure 1 further illustrates the addition of a second aqueous dispersion 40 including a mixture of two varieties of polymeric latex particles 42 to the slurry 30 of dispersed titanium dioxide particles 32 to form initially a formulated coating composition 50 including a mixture of dispersed titanium dioxide particles 32 and the two varieties of polymeric latex particles 42. The mixture of polymeric latex particles 42 includes a first type of polymeric latex particle 44 selected to adsorb strongly

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to the dispersed titanium dioxide particles 32 to form composite particles 46 including both a titanium dioxide particle 46a and a plurality of adsorbed selected polymer latex particles 46b. In one embodiment, the selected polymeric latex particles 46b have an effective glass transition temperature which is at least about room temperature or about 20 °C, and more preferably at least about 50 °C. In addition, the mixture of polymeric latex particles 42 includes a second type of polymeric latex particle 48 having a relatively low effective glass transition temperature, such as below about 10 °C, and serving ultimately to form a binder for the the composite particles 46.

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In another preferred embodiment, both the selected polymeric latex particles 46b and the second type of polymeric latex particles 48 have relatively low effective glass transition temperatures, such as below about 10 °C, and both serve as binders for the titanium dioxide particles.

In addition, other components such as low molecular weight cosolvents, plasticizers, anti-fungal and anti-bacterial agents, stabilizers and the like can be added (not shown) to provide a fully formulated coating composition. The fully formulated coating composition is then applied by conventional means to a surface 62 and 25 permitted to dry and cure, thus forming a coating film 60 in which the composite particles 46 including the titanium dioxide pigment particles 12 are imbedded in a continuous polymeric film 64 formed from the second variety of polymeric latex particles 42. The spacing of the 30 titanium dioxide particles 12 in the film 60 is not random, but instead is improved in comparison with prior art coatings films, so as to provide enhanced opacity.

Figure 2 schematically illustrates the distribution of titanium dioxide particles in a coating film formed from an aqueous composition prepared using a conventional prior art

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process. After thorough mixing of coating compositions, particles tend to be distributed in a random manner through the film, with many particles not contributing as much to the total light scattering of the film because they are closer to other particles than optimal. In contrast, Figure 3 schematically represents the distribution of titanium dioxide particles in a coating film formed from an aqueous coating composition using the process of the present invention. In this case, the particles are more regularly distributed in the film, resulting in greater effective light scatter per particle. Alternatively, fewer particles will provide a desired degree of opacification, and the coating PVC can be lower than if a prior art titanium dioxide dispersion method were used.

In the process of the present invention, the dispersed inorganic material particles can have negative surface potentials and charges, the negative charges serving to stabilize the dispersed inorganic material particles against homoaggregation. Similarly, the selected polymeric latex particles can have negative surface potentials and surface charges, also serving to stabilize the individual polymeric latex particles against homoaggregation or homocoagulation. However, despite the mutually repulsive electrical forces exerted on either type of particle, the selected polymeric latex particles are found to adsorb strongly, and even irreversibly, to the surface of the dispersed inorganic material particles. The mechanism for the adsorption process is not well understood. It is believed that the polyelectrolyte adsorbed to the surface of the inorganic material particles may be in equilibrium with polyelectrolyte dispersant dissolved in the aqueous medium, and that the selected polymeric latex particles may compete with the polyelectrolyte dispersant for binding sites on the surface of the inorganic material particles. However, it is known that adsorption of the selected polymeric latex particles may require several days after preparing the

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mixture including both the dispersed inorganic material particles and the selected polymeric latex particles. It is believed that adsorption of the selected polymeric latex particles is accompanied by reduction in the amount of adsorbed polyelectrolyte dispersant on the inorganic material particles, or a net displacement of the polyelectrolyte dispersant from the inorganic material particles.

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An alternative way to stabilize colloidal particles against uncontrolled aggregation is to provide a surface 10 layer or sheath of water soluble polymer around the colloidal particle, the chains of which polymer extend into the aqueous phase. This provides a steric barrier between the particles and between such a particle and other surfaces. A variety of water soluble polymers are suitable, 15 e.g., polyacrylamide, polyvinyl alcohol, polyethylene oxide, water soluble polymers derived from cellulose. The water soluble polymers may be suitably attached to the particle surfaces by several means, among them by chemical grafting, through polymerizable unsaturated groups linked to the water soluble polymer, as well as by physical adsorption, often enhanced by the presence of a hydrophobic group attached to the water soluble polymer. The hydrophobic group may be simply a hydrocarbon chain or a water insoluble block polymer attached to the water soluble 25 portion of the polymer. Steric stabilization has been thoroughly described by a number of authors, e.g., D.H. Napper, Polymeric Stabilization of Colloidal Dispersions, Academic Press, 1983; E.D. Goddard and B. Vincent, Polymer Adsorption and Dispersion Stability, ACS Symposium Series 30 240, 1984. Partial to thorough aggregation of sterically stabilized particles may be accomplished by reducing the solvency of the medium for the attached polymer chains, e.g., by variations in temperature, pressure, or composition of the liquid medium as described by Napper, 1983, chapter 5. For instance, a dispersion sterically stabilized with

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polyethylene oxide chains can be destabilized by the addition of salt or a water miscible nonsolvent for the polymer.

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Thus, in the process of the present invention, sterically stabilized polymeric latex particles can adsorb onto an inorganic material particle surface to provide composite particles. Hence, the present invention provides a process for preparing an aqueous dispersion of composite particles, the process comprising suspending inorganic material particles such as titanium dioxide particles in an 10 aqueous medium. In this aspect, the process further includes suspending the selected polymeric latex particles in the aqueous medium, the selected polymeric latex particles being stabilized against homoaggregation by polymeric steric stabilization. Also included is mixing the 15 aqueous medium containing the inorganic material particles and the polymeric latex particles, the polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles.

In the present invention the polymeric latex particles employed are polymerized in the absence of inorganic material particles, as distinguished from <u>in situ</u> processes.

Conditions which give gross heterocoagulation (i.e. flocculation) of the inorganic material particles and the polymeric latex particles can be avoided by providing both the inorganic material particles and the polymeric latex particles with negative surface potentials.

The polymeric latex particles can be prepared by a selected emulsion polymerization process which provides polymeric latex particles which nevertheless strongly adsorb to the inorganic material particles under these conditions, thus giving adsorption in a controlled manner to form the composite particles.

In some embodiments of the present invention inorganic material particles are added to an aqueous dispersion of polymeric latex particles. In particular, in these

embodiments the present invention provides a process for preparing an aqueous dispersion of composite particles, the composite particles each comprising a plurality of selected polymeric latex particles adsorbed onto an inorganic material particle. In these embodiments the process comprises:

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- a) preparing a dispersion or suspension of polymeric latex particles in an aqueous medium, the polymeric latex particles having a negative surface potential and being prepared by a preparative emulsion polymerization process in the absence of inorganic material particles, the preparative emulsion polymerization process being selected from the class consisting of:
- 1) emulsion polymerization processes employing a
 15 monomer mixture including at least one polymerizable
 ethylenically unsaturated acid-functional monomer, the at
 least one polymerizable ethylenically unsaturated acidfunctional monomer being selected from the class consisting
 of:

20 A) itaconic acid; and

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- B) dihydrogen phosphate esters of an alcohol, the alcohol containing a polymerizable olefinic group;
- 2) emulsion polymerization processes preferentially providing acid-functionality proximate the surface of the polymeric latex particle; and
- 3) emulsion polymerization processes employing a monomer mixture including at least one ethylenically unsaturated acid-functional monomer, the total weight of ethylenically unsaturated acid-functional monomer comprising at least about ten percent by weight of the polymeric latex solids;
- b) suspending inorganic material particles in the aqueous medium, the inorganic material particles having a negative surface potential in the aqueous medium, the negative surface potential of the inorganic material particles and the polymeric latex particles stabilizing the

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inorganic material particles and the polymeric latex particles against gross heterocoagulation, the polymeric latex particles being selected, by the use of the preparative emulsion polymerization process ,to adsorb onto the inorganic material particles in a controlled manner, and

c) mixing the aqueous medium containing the inorganic material particles and the selected polymeric latex particles, the selected polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles.

More specifically, in this set of embodiments, the present invention provides a process for preparing an aqueous dispersion of composite particles, the composite particles each comprising a plurality of selected polymeric latex particles adsorbed onto an inorganic material particle, the process comprising

- a) preparing a dispersion or suspension of selected polymeric latex particles in an aqueous medium, the polymeric latex particles having a negative surface potential and being prepared by a preparative emulsion polymerization process in the absence of inorganic material particles, the preparative emulsion polymerization process employing a monomer mixture including at least one polymerizable ethylenically unsaturated acid-functional monomer selected from the class consisting of monobasic and polybasic monomers having acidic functional groups including at least one proton with an acid equilibrium constant (pKa) \$of from about 4 to 8;
- aqueous medium, the inorganic material particles in the aqueous medium, the inorganic material particles having a negative surface potential in the aqueous medium, the negative surface potential of the inorganic material particles and the polymeric latex particles stabilizing the inorganic material particles and the polymeric latex

 35 particles against gross heterocoagulation, the polymeric latex particles being selected, by the use of the

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preparative emulsion polymerization process, to adsorb onto the inorganic material particles in a controlled manner, and

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c) mixing the aqueous medium containing the inorganic material particles and the selected polymeric latex particles, the selected polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles.

For example, the aqueous dispersion of composite particles can be prepared according to the process of these embodiments, such that the composite particles each comprise a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including polymerized residues of itaconic acid.

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Similarly, the present invention provides a process for preparing an aqueous composition, such as a pigment 15 slurry, or a coating composition, including such dispersed composite particles. For example, the pigment slurry can comprise an aqueous dispersion of composite particles prepared according to the process of these embodiments, with 20 the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including dihydrogen phosphate ester functional groups. Similarly, the pigment slurry can comprise an aqueous dispersion of composite particles prepared according to the process of these 25 embodiments, wherein the composite particles each comprise a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, and the polymeric latex particles include polymerized residues of itaconic acid. Further, in a coating composition prepared according to the process of 30 these embodiments, the coating composition can comprise an aqueous dispersion of composite particles, the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including polymerized residues of 35 itaconic acid.

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In another set of embodiments a separate aqueous suspension of the inorganic material particles is prepared and mixed with the aqueous dispersion of the polymeric latex particles. In this set of embodiments, the present invention provides process for preparing an aqueous dispersion of composite particles or aqueous composition including such dispersed composite particles, the composite particles each comprising a plurality of selected polymeric latex particles adsorbed onto an inorganic material particle, the process comprising:

- a) preparing a dispersion or suspension of polymeric latex particles in a first aqueous medium, the polymeric latex particles being prepared by a preparative emulsion polymerization process in the absence of inorganic material particles, the preparative emulsion polymerization process being selected from the class consisting of:
- 1) emulsion polymerization processes employing a monomer mixture including at least one polymerizable ethylenically unsaturated acid-functional monomer, the at least one polymerizable ethylenically unsaturated acid-functional monomer being selected from the class consisting of:
 - A) itaconic acid; and
- B) dihydrogen phosphate esters of an alcohol, the alcohol containing a polymerizable olefinic group;
 - 2) emulsion polymerization processes preferentially providing acid-functionality proximate the surface of the polymeric latex particle; and
- 3) emulsion polymerization processes employing a monomer mixture including at least one ethylenically unsaturated acid-functional monomer, the total weight of ethylenically unsaturated acid-functional monomer comprising at least about ten percent by weight of the polymeric latex solids;
- b) suspending inorganic material particles in a second aqueous medium; and

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c) mixing the first and second aqueous media containing the polymeric latex particles and the inorganic material particles respectively, the polymeric latex particles and the inorganic material particles each having a negative surface potential in the mixed media, the negative surface potential of the inorganic material particles and the polymeric latex particles stabilizing the inorganic material particles and the polymeric latex particles against gross heterocoagulation, the polymeric latex particles being selected, by the use of the preparative emulsion polymerization process, to adsorb onto the inorganic material particles in a controlled manner, the polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles. --

More specifically, in this second set of embodiments, the present invention provides a process for preparing an aqueous dispersion of composite particles, the composite particles each comprising a plurality of selected polymeric latex particles adsorbed onto an inorganic material particle, the process comprising:

- a) preparing a dispersion or suspension of polymeric latex particles in an aqueous medium, the polymeric latex particles having a surface potential and being prepared by a preparative emulsion polymerization process in the absence of inorganic material particles, the preparative emulsion polymerization process being selected from the class consisting of:
- 1) emulsion polymerization processes employing a monomer mixture including at least one polymerizable ethylenically unsaturated acid-functional monomer, the at least one polymerizable ethylenically unsaturated acid-functional monomer being selected from the class consisting of:
 - A) itaconic acid; and
- B) dihydrogen phosphate esters of an alcohol, the alcohol containing a polymerizable olefinic group;

2) emulsion polymerization processes preferentially providing acid-functionality proximate the surface of the polymeric latex particle; and

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- 3) emulsion polymerization processes employing a monomer mixture including at least one ethylenically unsaturated acid-functional monomer, the total weight of ethylenically unsaturated acid-functional monomer comprising at least about ten percent by weight of the polymeric latex solids;
- b) suspending inorganic material particles in the aqueous medium, the inorganic material particles having a surface potential in the aqueous medium of the same sign as the surface potential of the polymeric latex particles, the surface potential of the inorganic material particles and the polymeric latex particles stabilizing the inorganic material particles and the polymeric latex particles against gross heterocoagulation, the polymeric latex particles being selected, by the use of the preparative emulsion polymerization process, to adsorb onto the inorganic material particles in a controlled manner, and
 - c) mixing the aqueous medium containing the inorganic material particles and the selected polymeric latex particles, the selected polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles.

Alternatively, in this second set of embodiments, the present invention provides a process for preparing an aqueous dispersion of composite particles, the composite particles each comprising a plurality of selected polymeric latex particles adsorbed onto an inorganic material particle, which can be expressed as follows: The process comprises:

a) preparing a dispersion or suspension of polymeric latex particles in a first aqueous medium, the polymeric latex particles being prepared by a preparative emulsion polymerization process in the absence of inorganic

material particles, the preparative emulsion polymerization process being selected from the class consisting of:

- nonomer mixture including at least one polymerizable ethylenically unsaturated acid-functional monomer, the at least one polymerizable ethylenically unsaturated acid-functional monomer being selected from the class consisting of:
 - A) itaconic acid; and
- B) dihydrogen phosphate esters of an alcohol, the alcohol containing a polymerizable olefinic group;
 - 2) emulsion polymerization processes preferentially providing acid-functionality proximate the surface of the polymeric latex particle; and
- 3) emulsion polymerization processes employing a monomer mixture including at least one ethylenically unsaturated acid-functional monomer, the total weight of ethylenically unsaturated acid-functional monomer comprising at least about ten percent by weight of the polymeric latex solids;
 - b) suspending inorganic material particles in a second aqueous medium; and

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c) mixing the first and second aqueous media containing the polymeric latex particles and the inorganic material particles respectively, the polymeric latex particles and the inorganic material particles each having a surface potential of the same sign in the mixed media, the surface potential of the inorganic material particles and the polymeric latex particles stabilizing the inorganic material particles and the polymeric latex particles against gross heterocoagulation, the polymeric latex particles being selected, by the use of the preparative emulsion polymerization process, to adsorb onto the inorganic material particles in a controlled manner, the polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles. --

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A third variation, in which previously dried latex particles are suspended in an aqueous suspension of inorganic material particles, is considered an obvious equivalent of these two sets of embodiments. These two sets of embodiments are considered to be mere variations in the sequence of steps. In some embodiments the surface potential of polymeric latex particles and the inorganic material particles have the same sign, but that sign may be either positive or negative.

The controlled adsorption of the polymeric latex 10 particles of the present invention is paradoxical and unexpected. High surface potential on like sign on both pigment and polymer particles is commonly understood to provide great stability through repulsive electrical forces. Similarly, a high concentration of surface acid, such as 15 thought to be provided by some of the emulsion polymerization processes which can be employed in preparing the selected polymeric latex particles for the present invention, might well be understood to provide for a high level of stabilizing repulsive forces with respect to 20 similarly charged pigment particles. Nevertheless, it has been found that selected polymeric latex particles thought to possess such high surface acid can adsorb in a controlled manner to such pigment particles. This phenomenon has been employed in the present invention to achieve important and 25 unexpected results, such as enhanced opacity, improved mechanical properties, and the like.

The surface characteristics of a particle suspended in an aqueous medium can often be adjusted by changing the characteristics of the aqueous medium. For example, many inorganic particles, including titanium dioxide particles, have acidic and/or basic functional groups at their surfaces, or can be treated to provide such groups at the particles' surfaces, such as by adsorption of anionic pigment dispersants and/or anionic surfactants, and the surface charge of these particles can be adjusted by varying

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the pH of the aqueous medium.

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Similarly, polymeric latex particles can be synthesized with acidic and/or basic functional groups at their surfaces, or can be treated to provide such groups at the particles' surfaces, such as by adsorption of anionic pigment dispersants and/or anionic surfactants, and the surface charge density of these particles can also be adjusted by varying the pH of the aqueous medium. case of synthetic polymeric latex particles, the surface charge density can also be controlled by adjusting the density of surface acid and/or base groups through selection of polymer composition and polymerization process variables. An emulsion polymerization process which provides a preferential distribution of acid functionality for the particles, such as proximate the surface of the particles, can be used to prepare the selected polymeric latex particles employed in one presently preferred embodiment of the present invention.

The entire accessible pH scale is useful in the practice of this invention although it is preferred for reasons of excessive particle solubility, corrosion both to substrates and to the skin, and the like, to restrict the range to approximately 2 to 12 and more preferably in the range of from about 4 to about 10.

Inorganic material particles which have been subjected to prior surface treatment processes can be used in the process of the present invention. For example, titanium dioxide is available with various types of prior surface treatments which provide surface coatings of alumina or silica or mixtures of both, the treated inorganic material particles each exhibiting differing adsorption characteristics, as discussed in A. Brisson et al., J. Coatings Tech. 63 59-65 (1991).

The adsorption process of the present invention is 35 believed to be dependent to some extent on the particle sizes of the polymeric latex particles and the inorganic

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material particles to be adsorbed thereto, as well as the amount of the polymeric latex and inorganic material particles in the aqueous adsorption medium. At a sufficiently high concentration, it can be postulated that the selected polymeric latex can adsorb onto and saturate the entire surface of the inorganic material particle's surface, and an empirical saturation level can be determined. It is believed that the empirical saturation level observed depends primarily on particle size, and that other factors such as surface charge, polymeric latex glass 10 transition temperature, and the like, have little effect. Preferably, enough of the selected polymeric latex is employed to obtain optimum performance properties in fully formulated products prepared using the process.

In preparing coating formulations possessing desirable 15 performance properties it is not necessary that the selected polymeric latex used in the adsorption process of the invention to also function as the polymeric binder for the coating or film, although this may be possible or even desirable in some cases. Instead, another polymeric latex 20 material, preferably having a glass transition temperature tending to provide good film formation properties under the application conditions, can be provided to serve this function. By "glass transition temperature" is meant the 25 second order phase transition temperature as determined empirically by mechanical methods as torsional braid analysis and the like or as calculated from monomer composition by the method of Fox. By "effective glass transition temperature" is meant the second order phase transition temperature of the polymeric material as modified 30 by the presence of low molecular weight species such as coalescents, solvent, and the like. The function of the selected polymeric latex in contrast is to adsorb to the inorganic material particles and to space them from one another in the film formed by the polymeric binder latex. To serve this function, it may be desirable that the

selected polymeric latex particles be relatively rigid, having a relatively high effective glass transition temperature under the application conditions, for example, at least about 20 °C, more preferably at least about 35 °C, and even more preferably at least about 50 °C. Thus, one presently preferred embodiment of the present invention provides an aqueous mixture of the selected, strongly adsorbing "hard" polymeric latex particles and the "soft", film-forming polymeric latex particles for use in preparing formulated aqueous compositions, such as coating compositions.

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With regard to the average particle size or diameter of the selected polymeric latex particles and the inorganic material particles employed in the present process, it is generally preferred that the selected polymeric latex particles have an average diameter of from about 20 nm to about four times that of the inorganic material particles, and more preferably from about 20 nm to about the same diameter as that of the inorganic material particles, especially in the case in which the inorganic material particles are titanium dioxide particles. In one presently preferred embodiment, in the case of selected, strongly adsorbing "hard" polymeric latex particles, an average polymer latex diameter from about 30 nm to about 100 nm is preferred, and an average polymer latex diameter from about 50 nm to 80 nm is especially preferred.

In a second presently preferred embodiment, in the case of selected, strongly adsorbing "soft" polymeric latex particles, an average polymeric latex particle diameter of from about 80 nm to 600 nm is preferred, and an average particle diameter of from about 100 nm to 400 nm is especially preferred.

However, the optimum relative particle sizes are strongly dependent on the rigidity of the polymeric latex particles, as measured, for example, by their glass transition temperature, and the pigment volume concentration

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of the coating or product formed from the composition including the composite particles. In at least some coating compositions, hiding is maximized when the polymeric latex particle diameter is greater than the titanium dioxide particle diameter.

The concentration of the titanium dioxide particles (and any other pigments which may be present in the composition) in a coating formulation is expressed in terms of the pigment volume concentration of the formulation. pigment volume concentration (hereinafter referred to as the "PVC") of a formulation is defined as the volume amount of inorganic particles, including titanium dioxide and other pigment particles as well as extender particles, present in the formulation, divided by the sum of the volume amount of such inorganic particles plus polymeric latex particle solids in the formulation. The overall concentration of pigment particles, extender particles and emulsion polymer particles in a formulation is typically expressed in terms of a percent volume solids for the formulation. The percent volume solids is an expression of the extent of the dilution of the solids in a liquid vehicle, such as water.

Through the selection of the polymeric latex composition and the nature of the adsorbed pigment dispersant, the adsorption of the selected polymeric latex particles on the surface of the inorganic material particle can occur substantially upon the intimate mixing of the selected polymeric latex with the inorganic material particles, or the adsorption process may extend over a substantial time, such as several days or weeks. As a practical matter, the fact that the adsorption process is not significantly completed until after an extended period of time is of little import, as most formulated compositions including both inorganic material particles and polymeric latex particles must be stored for an extended period between manufacture and application, to accommodate the requirements of the consumer of the products. For example,

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several months may elapse after preparation of an aqueous latex paint and its application, and the paint must be formulated so as to possess sufficient storage stability. Only in those instances in which the formulated aqueous composition must be used shortly after manufacture is . selection of the absorbing polymeric latex required to provide a high degree of adsorption on the inorganic material particles shortly after mixing.

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The polymeric latexes used in the practice of the present invention can have monomer compositions and particle sizes closely related to polymeric latex binders prepared by standard emulsion polymerization techniques known in the art. If it is otherwise desirable to employ conventional levels of a polyelectrolyte dispersant in a pigment dispersion, it may be possible to employ a strongly adsorbing polymeric latex to adsorb onto the inorganic material particles. Strongly adsorbing polymeric latex particles can be prepared by emulsion polymerization processes by selection of process variables and the monomer 20 composition.

In this invention, the polymeric latex particles are selected such that they can be used to adsorb to the inorganic material particles in the presence of a polyelectrolyte pigment dispersant already adsorbed to the surface of the inorganic material particles. As the polymeric latex particles are used in the presence of a pigment dispersant, it is preferable that the polymeric latex particles bind irreversibly to the inorganic material particles.

In a presently preferred embodiment of the process of 30 the present invention, the selected polymeric latex particles do not themselves provide a binder for the product formed by the fully formulated aqueous composition. Because they are not required to flow and form a film, the particles can be more rigid than those employed to provide a binder. 35 Thus, the polymeric particles can have a higher effective

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glass transition temperature than polymeric particles employed to serve as binder, and/or can include rigidifying levels of crosslinking. In this case the polymeric latex can also be prepared by standard emulsion polymerization techniques, but typically will not be suitable for use as a film-forming binder.

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The polymeric latex particles used in these embodiments are preferably selected based on an ability to adsorb rapidly and irreversibly onto the inorganic material particles. Such particles are not displaced from the surface of the inorganic material particles upon addition of a second variety of polymeric latex having a monomer composition and physical properties which differ from that of the initial polymeric latex.

This aspect of the invention advantageously provides enhanced flexibility in formulation. For example, the second variety of polymeric latex particles can be polymeric latex which has a monomer composition and physical properties which have been optimized for use as a binder for a specific coating application. Some of these optimized properties may not be consistent with use of this specific polymeric latex as an adsorbent in the process of the present invention.

In this aspect, dispersion of the inorganic material particles can be otherwise effected, as with a polyelectrolyte pigment dispersant, with a mixture of irreversibly adsorbing polymeric latex particles and binder latex particles being subsequently mixed with the dispersed inorganic material particles, the irreversibly adsorbing polymeric latex particles advantageously spacing the inorganic material particles in the product of the formulated composition, while the binder latex particles provide a continuous film otherwise optimized for the application. This is especially useful when titanium dioxide is employed as the inorganic material, and the improved spacing results in enhanced opacity and hiding.

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Strongly adsorbing polymeric latex particles useful in the process of this invention may be prepared by several alternative processes. In one such process, emulsion polymerization process variables are controlled to provide a preferred distribution of acid functionality for the polymer latex particles. In particular, processes which provide carboxylate functionality preferentially at the surface of the polymeric latex particles are preferred. In another process, the polymerization process is employed in which the total weight of ethylenically unsaturated acid-functional 10 monomer comprises at least about five percent by weight of the polymeric latex solids, and preferably at least about ten percent by weight. In still another process for preparing strongly adsorbed polymeric latex particles, the monomer mixture from which the polymeric latex is 15 polymerized includes at least one dihydrogen phosphate ester of an alcohol in which the alcohol contains a polymerizable olefinic group, such as a polymerizable vinyl group. Examples of such dihydrogen phosphate esters include allyl 20 phosphate, allyl cellosolve phosphate, hydroxyethyl methacrylate phosphate, the mono- or diphosphate of bis(hydroxyethyl) fumarate or itaconate, etc. In particular, polymeric latex particles polymerized from monomer mixtures including the dihydrogen phosphate ester of 2-hydroxyethyl methacrylate are preferred. In another such 25 process, the monomer mixture from which the selected polymeric latex is polymerized includes itaconic acid as a polymerizable, ethylenically unsaturated acid-functional monomer.

without intending to be bound by any particular explanation of the mechanism of the process of the present invention, it is presently believed that selected polymeric latexes having acidic functionality with a pKa intermediate between weak and strong acids can be effective in the process of the present invention, such as those polymerized from monomer including copolymerizable, ethylenically

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unsaturated monomers with acidic functional groups with a pKa from about 4 to 8, more preferably about 6. It should be noted that the pKa of an acidic functional group can vary depending upon its enviorment. For example, the measured pKa of an acid functional group can change when a polymerizable monomer bearing that group is copolymerized with more hydrophobic comonomers and the environment of the acid functional group becomes more hydrophobic. It is theorized that under basic conditions, such as are frequently employed in aqueous polymeric latex-based coating compositions, the ionic conjugate species of these acidic functional groups enhance the adsorption of the selected polymeric latex particles on the inorganic material particles to provide the composite particles.

In general it is believed that reducing the effective glass transition temperature of the polymeric latex (for example, by changing the monomer composition) increases the tendency to adsorb strongly, as does increasing the level of acid-functional monomers, and in particular, the level of phosphate ester-functional monomers or itaconic acid, in the composition, or enhancing the surface concentration of acid-functional monomers. However, it is often desirable to employ a relatively hard, strongly adsorbing polymeric latex, to maximize the improvements provided by the present invention in the performance properties of aqueous composition including the composite particles.

Thus, the present invention provides aqueous dispersions of composite particles, as well as coating compositions and pigment slurries including such composite particles, wherein the composite particles each comprise a plurality of polymeric latex particles adsorbed to an inorganic material particle, the polymeric latex particles including, for example, dihydrogen phosphate ester functional groups.

These dihydrogen phosphate ester functional groups are believed to result from the polymeric latex particles being

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polymerizable dihydrogen phosphate-functional monomer. The polymerizable dihydrogen phosphate-functional monomer can be selected from the dihydrogen phosphate esters of an alcohol, the alcohol including a polymerizable group selected from polymerizable vinyl groups and polymerizable non-vinyl olefinic groups. The polymerizable dihydrogen phosphate-functional monomer can be selected from the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate, and mixtures of the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate and the phosphoric acid diester of 2-hydroxyethyl methacrylate.

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Similarly, the present invention provides aqueous dispersions of composite particles, and coating compositions and pigment slurries including such composite particles, in which the composite particles each comprising a plurality of polymeric latex particles adsorbed to an inorganic material particle, the polymeric latex particles including polymerized residues of itaconic acid.

The polymeric latex used in the present invention can be prepared by any technique known in the art, such as suspension polymerization or emulsion polymerization. Emulsion polymerization techniques for preparing aqueous dispersions of latex polymer particles from ethylenically unsaturated monomers are well known in the polymer arts, and any conventional emulsion technique can be used, such as single and multiple shot batch processes, and continuous processes. If desired, a monomer mixture can be prepared and added gradually to the polymerization vessel. The monomer composition within the polymerization vessel can be varied during the course of the polymerization, such as by altering the composition of the monomer being fed into the vessel. Both single and multiple stage polymerization techniques can be used. The latex polymer particles can be prepared using a seed polymer emulsion to control the number of particles produced by the emulsion polymerization as is

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known in the art. The particle size of the latex polymer particles can be controlled by adjusting the initial surfactant charge as is known in the art. The preparation of polymeric latexes is discussed generally in D.C.

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Blackley, <u>Emulsion Polymerization</u> (Wiley, New York, 1975).

The preparation of acrylic polymeric latexes is described in, for example, <u>Emulsion Polymerization of Acrylic Polymers</u>, Bulletin, Rohm and Haas Company, Philadelphia,

A polymerization initiator can be used in carrying out
the polymerization of the polymeric latex particles.

Examples of polymerization initiators which can be employed include polymerization initiators which thermally decompose at the polymerization temperature to generate free radicals. Examples include both water-soluble and water-insoluble

15 species. Examples of free radical-generating initiators which can be used include persulfates, such as ammonium or alkali metal (potassium, sodium or lithium) persulfate; azo compounds such as 2,2'-azo-bis(isobutyronitrile),

2,2'-azo-bis(2,4-dimethylvaleronitrile), and

1-t-butyl-azocyanocyclohexane); hydroperoxides such as t-butyl hydroperoxide and cumene hydroperoxide; peroxides such as benzoyl peroxide, caprylyl peroxide, di-t-butyl peroxide, ethyl 3,3'-di(t-butylperoxy) butyrate, ethyl 3,3'-di(t-amylperoxy) butyrate, t-amylperoxy-2-ethyl

hexanoate, and t-butylperoxy pivilate; peresters such as t-butyl peracetate, t-butyl perphthalate, and t-butyl perbenzoate; as well as percarbonates, such as di(1-cyano-1-methylethyl)peroxy dicarbonate; perphosphates, and the like.

Polymerization initiators can be used alone or as the oxidizing component of a redox system, which also includes a reducing component such as ascorbic acid, malic acid, glycolic acid, oxalic acid, lactic acid, thiogycolic acid, or an alkali metal sulfite, more specifically a

hydrosulfite, hyposulfite or metabisulfite, such as sodium hydrosulfite, potassium hyposulfite and potassium

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metabisulfite, or sodium formaldehyde sulfoxylate. The reducing component is frequently referred to as an accelerator.

The initiator and accelerator, commonly referred to as catalyst, catalyst system or redox system, can be used in proportion from about 0.001% to 5% each, based on the weight of monomers to be copolymerized. Accelerators such as chloride and sulfate salts of cobalt, iron, nickel or copper can be used in small amounts. Examples of redox catalyst systems include tertbutyl hydroperoxide/sodium formaldehyde 10 sulfoxylate/Fe(II), and ammonium persulfate/sodium bisulfite/sodium hydrosulfite/Fe(II). The polymerization temperature can be from room temperature to about 90°C, and can be optimized for the catalyst system employed, as is conventional.

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Chain transfer agents can be used to control polymer molecular weight, if desired. Examples of chain transfer agents include mercaptans, polymercaptans and polyhalogen compounds. Examples of chain transfer agents which may be 20 used include alkyl mercaptans such as ethyl mercaptan, n-propyl mercaptan, n-butyl mercaptan, isobutyl mercaptan, t-butyl mercaptan, n-amyl mercaptan, isoamyl mercaptan, t-amyl mercaptan, n-hexyl mercaptan, cyclohexyl mercaptan, n-octyl mercaptan, n-decyl mercaptan, n-dodecyl mercaptan; alcohols such as isopropanol, isobutanol, lauryl alcohol and t-octyl alcohol; halogenated compounds such as carbon tetrachloride, tetrachloroethylene, and tricholorobromoethane. Generally from 0 to 10% by weight, based on the weight of the monomer mixture, can be used. The polymer molecular weight can be controlled by other techniques known in the art, such as selecting the ratio of initiator to monomer.

Catalyst and/or chain transfer agent can be dissolved or dispersed in separate or the same fluid medium and gradually added to the polymerization vessel. Monomer, either neat or dissolved or dispersed in a fluid medium, can

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be added simultaneously with the catalyst and/or the chain transfer agent. Amounts of initiator or catalyst can be added to the polymerization mixture to "chase" residual monomer after polymerization has been substantially completed to polymerize the residual monomer as is well known in the polymerization arts.

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Aggregation of polymeric latex particles is typically discouraged by including a stabilizing surfactant in the polymerization mix. In general, the growing latex particles are stabilized during emulsion polymerization by one or more 10 surfactants such as an anionic or nonionic surfactant, or a mixture thereof, as is well known in the polymerization art. Many examples of surfactants suitable for emulsion polymerization are given in McCutcheon's Detergents and Emulsifiers (MC Publishing Co., Glen Rock, NJ), published 15 annually. Other types of stabilizing agents, such as protective colloids, can also be used. However, it is preferred that the amount and type of stabilizing surfactant or other type of stabilizing agent employed during the polymerization reaction be selected so that residual 20 stabilizing agent in the aqueous product of the polymerization reaction does not significantly interfere with the subsequent adsorption of the polymeric latex to the inorganic material. Further, charged initiator fragments 25 and copolymerized monomer bearing charged functional groups such as copolymerized acid-functional monomers are known to contribute to the stability of the resulting polymeric latex particles. It is important to note that stabilizing surfactants, copolymerized acidic monomers, residual initiator fragments, and the like, which tend to 30 contribute to stability of the polymeric latex particles with respect to homoaggregation or homocoagulation, may also tend to stabilize the polymeric latex particles with respect to heterocoagulation or heteroaggregation in general, and composite particle formation in particular. Thus adjusting 35 polymerization conditions to provide a desired level of

residual initiator fragments and surface acid may be very important in providing polymeric latex particles for use in the controlled adsorption process of the present invention.

The polymeric latexes useful in the process of the

present invention can be prepared from a wide range of
polymerizable monomers, such as, for example,
monoethylenically unsaturated monomers, including alpha,
beta-monoethylenically unsaturated monomers such as alkyl
acrylates and methacrylates. By "acrylic polymeric latex"

is meant a polymeric latex polymerized from monomers
comprising substantially polymerizable monomers including
the acryl group (-COCH=CH₂) or methacryl (-COC(CH₃)=CH₂)
group, and specifically greater than about 80 weight percent
(meth)acrylic monomers, based on the total monomer weight.

Mixtures of polymeric latexes can also be used.

Examples of polymeric latexes which can be employed in the process of the present invention include those polymerized from ethylenically unsaturated monomers, such as alpha, beta-ethylenically unsaturated monomers, including styrene, butadiene, alpha-methylstyrene, vinyltoluene, vinylnaphthalene, ethylene, vinyl acetate, vinyl versatate, vinyl chloride, vinylidene chloride, acrylonitrile, methacrylonitrile, (meth)acrylamide, various (C1-C20)alkyl esters of (meth)acrylic acid; for example, methyl (meth)acrylate, ethyl (meth)acrylate, n-butyl (meth)-

25 (meth)acrylate, ethyl (meth)acrylate, n-butyl (meth) acrylate, isobutyl (meth)acrylate, 2-ethylhexyl
(meth)acrylate, cyclohexyl (meth)acrylate, n-octyl
(meth)acrylate, n-decyl (meth)acrylate, n-dodecyl
(meth)acrylate, tetradecyl (meth)acrylate, n-amyl

(meth)acrylate, neopentyl (meth)acrylate, cyclopentyl (meth)acrylate, lauryl (meth)acrylate, oleyl (meth)acrylate, palmityl (meth)acrylate, and stearyl (meth)acrylate; other (meth)acrylates such as isobornyl (meth)acrylate, benzyl (meth)acrylate, phenyl (meth)acrylate, 2-bromoethyl

(meth)acrylate, 2-phenylethyl (meth)acrylate, and 1-naphthyl (meth)acrylate; alkoxyalkyl (meth)acrylate such as

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ethoxyethyl (meth) acrylate; mono-, di-, and trialkyl esters of ethylenically unsaturated di- and tricarboxylic acids and anhydrides, such as ethyl maleate, dimethyl fumarate, trimethyl aconitate, and ethyl methyl itaconate. As used in the present specification and claims, "(meth) acrylate" denotes both "acrylate" and "methacrylate" and "acrylic."

The ethylenically unsaturated monomer can also include at least one multi-ethylenically unsaturated monomer effective to raise the molecular weight and crosslink the polymer. Examples of multi-ethylenically unsaturated monomers that can be used include allyl (meth)acrylate, tripropylene glycol di(meth)acrylate, diethylene glycol di(meth)acrylate, ethylene glycol di(meth)acrylate, 1,6-hexanediol di(meth)acrylate, 1,3-butylene glycol (meth)acrylate, polyalkylene glycol di(meth)acrylate, diallyl phthalate, trimethylolpropane tri(meth)acrylate, divinylbenzene, divinyltoluene, trivinylbenzene, and divinylnaphthalene.

In addition to those monomers which are "functional" in the sense of including one or more polymerizable ethylenically unsaturated groups, monomers which also include one or more additional functional groups can be used in preparing the polymeric latexes used in the process of this invention. An important class of these functional monomers is that made up of those polymerizable ethylenically unsaturated monomers having acidic functional groups. Examples of these include acrylic acid, methacrylic acid, itaconic acid, beta-acryloxypropionic acid and higher monoacidic oligomers of acrylic acid, ethacrylic acid, alpha-chloroacetic acid, alpha-vinylacrylic acid, crotonic acid, alpha-phenylacrylic acid, cinnamic acid, chlorocinnamic acid, beta-styrylacrylic acid, itaconic acid, maleic acid, dihydrogen phosphate esters of an alcohol in which the alcohol also contains a polymerizable vinyl or olefinic group, such as allyl phosphate, allyl Cellosolve

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phosphate, mono- or diphosphate of bis(hydroxy-methyl) fumarate or itaconate, derivatives of (meth)acrylic acid esters, such as, for example, phosphates of hydroxyalkyl(meth)acrylates including 2-hydroxyethyl (meth)acrylate, such as Kayamer (trademark of Nihon Kayaku Co., Ltd.) PM-1, PM-2, PA-1, and PA-2 monomers, 3-hydroxypropyl (meth)acrylates, and the like. Examples of dihydrogen phosphate ester derivatives which can be employed include vinylbenzyl phosphates, and:

 $CH_2 = CCH_3CO_2CH_2CH_2OPO (OH)_2$ $CH_2 = CCH_3CO_2CH_2CH [OPO (OH)_2] CH_3$ $CH_2 = CCH_3CO_2CH_2CH_2CH_2OPO (OH)_2$ $CH_2 = CCH_3CO_2CH_2CHOHCH_2OPO (OH)_2$

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Thus, the present invention also provides aqueous
dispersions of composite particles, as well as coating
compositions and pigment slurries including such composite
particles, in which the composite particles include
polymeric latex particles polymerized from monomer including
at least one polymerizable dihydrogen phosphate-functional
monomer selected from the dihydrogen vinylbenzyl phosphates,
and CH₂=CCH₃CO₂CH₂CH₂OPO(OH)₂, CH₂=CCH₃CO₂CH₂CH[OPO(OH)₂] CH₃,
CH₂=CCH₃CO₂CH₂CH₂CH₂OPO(OH)₂, and CH₂=CCH₃CO₂CH₂CHOHCH₂OPO(OH)₂.

Small amounts of acid-functional copolymerizable monomer, such as methacrylic acid and/or acrylic acid, are typically included in preparing polymeric latexes to confer colloidal stability. As noted above, in one embodiment of the present invention, preparation of polymeric latex particles from monomer mixture including itaconic acid is preferred. In another preferred embodiment, a relatively large amount of acid-functional copolymerizable monomer, such as methacrylic acid, for example, at least about five percent by weight of total polymer solids, and preferably at least about ten percent by weight of total polymer solids, is included in the monomer mix from which the polymeric latex particles are prepared. In yet another presently preferred embodiment, the polymerization is carried out so

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as to preferentially provide acid-functionality proximate the surface of the polymeric latex particles. In another presently preferred embodiment of the present invention, as noted above, the polymeric latex employed is polymerized from monomer mixture including at least one phosphoric acid partial ester derivative of a (meth) acrylic acid ester, preferably including a dihydrogen phosphate monoester.

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In the emulsion copolymerization of ethylenically unsaturated monomers including minor amounts (such as less than about ten percent by weight) of ethylenically unsaturated carboxylic acid-functional comonomers such as (meth)acrylic acid, the resulting polymeric acid groups may be buried inside the colloidal polymeric particles produced by the polymerization and consequently inaccessible to species dissolved in the aqueous phase, or the polymeric acid groups may be at or near the surface of the particles, or even present in the aqueous phase in the form of water-soluble polymer. In addition, some of the acidfunctional monomer may remain unpolymerized. This residual monomer is ususally to be found in the aqueous phase. functionality which is at or near the surface of the particles (that is, proximate the surface of the polymeric latex particles) can be measured by potentiometric or conductometric titration techniques.

For example, if the polymeric latex particles are treated with ion-exchange resin and then titrated conductometrically, the amount of carboxylic acid at or near the surface of the particles can be measured as described in H.J. Van den Hul and J. W. Vanderhoff, Electro. Chem.

Interfac. Electrochem., 37 161-182 (1972). High levels of surface carboxylic acid promote the adsorption process of this invention.

For the same mixture of monomers, the amount of surface carboxylic acid groups in the resulting polymeric latex particles depends on the specific emulsion polymerization employed. A number of factors can be

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important, including the particle size of the resulting polymeric latex, the pH of the polymerization medium, how the monomer is added to the polymerization reactor, and the choice of carboxylic acid monomer. For instance, if a mixture of 50 percent by weight butyl acrylate, 49 percent methyl methacrylate, and one percent methacrylic acid is gradually added to a polymerization reactor after the polymerization has been initiated, such as in a conventional gradual addition process, to give 200 nm polymer particles, 10 titration indicates that about 25 percent of the total methacrylic acid is "surface acid," at or near the surface of the polymeric latex particles. If acrylic acid is substituted for methacrylic acid, the surface acid can increase. Similarly, if the process is altered so that all the monomer is added to the reactor, and the polymerization 15 is then initiated, as is practical in a small-scale batch process, the surface acid can be found to be somewhat higher. In the present application, a process which preferentially provides acid functionality proximate the 20 surface of the polymer latex particles is one which provides greater surface acid functionality than a conventional gradual addition process, or a conventional "single shot" batch process (such as a conventional process in which base is not added to the polymerization mix and the acidic functional groups consequently remain substantially 25 nonionized through the polymerization process), as determined by potentiometric titration.

There are a number of ways known in the art for increasing the fraction of the acid that becomes surface acid. For example, if enough base is added during the polymerization to partially neutralize the carboxylic acid, the surface acid can be increased measurably. However, higher levels of base can decrease surface acid. If the carboxylic acid monomer is added to the reactor in a nonuniform manner, the amount of surface acid can be increased. Examples of nonuniform carboxylic acid monomer

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additions that increase surface acid are described in K.L. Hoy, <u>J. Coat. Tech.</u>, 51 27-41 (1979).

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Other types of copolymerizable functional monomers can also be included, depending on the ultimate application for which the product produced by the process of the present invention is intended. For example, small amounts of adhesion-promoting copolymerizable monomers can also be included. Examples of other types of functional monomers include hydroxy-functional monomers such as 2-hydroxethyl (meth)acrylate, amino-functional monomers such as dimethylaminoethyl (meth)acrylate, epoxy-functional monomers, such as glycidyl methacrylates and the like.

Examples of synthetic polymeric latexes which can be used include acrylic copolymer latexes, such as butyl acrylate/methyl methacrylate/acid and ethyl acrylate/methyl methacrylate/acid copolymer latexes; vinyl acetate homopolymer and copolymer latexes, including ethylene/vinyl acetate copolymer latexes, styrene/butadiene copolymer latexes, and the like.

By selecting the type and concentration of monomers having known glass transition temperatures and acid functionalities, polymeric latexes having properties useful in the process of the invention can be prepared.

The particle size, particle size distribution, and molecular weight of the polymeric latex can be selected by the choice of a particular polymerization process, as for example through the use of particular initiator and chain transfer systems, as is well known in the polymerization arts. The average particle size and the molecular weight of the polymeric latex are important with regard to both the adsorption onto the inorganic material particles, as well as with respect to the properties of the polymeric latex when it is employed as a binder in the the fully formulated aqueous composition including the composite particles.

Preferably, the polymeric latex used in the process of the present invention of adsorbing onto the inorganic

material particles has an average particle diameter in the range of from approximately four times the average particle diameter of the inorganic material particles down to about 20 nm. For example, if titanium dioxide particles are to be used, and those particles have an average particle size of about 200 nanometers, the polymeric latex particle should have an average particle size in the range of from about 20 to less than about 800 nanometers. By "average particle size" or "average particle diameter" is meant an average determined experimentally by the quasielastic light scattering technique, such as provided, for example, by the Model BI-90 Particle Sizer, of Brookhaven Instruments Corp.

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The inorganic material particles, such as titanium dioxide particles, can be pretreated to provide inorganic surface coatings, such as coatings of silica, aluminum oxide, or mixtures of silica and aluminum oxide, on titanium dioxide particles. In addition, small molecular species, such as silanes or titanates, can be adsorbed on and reacted with surface of the titanium dioxide particles, and the resulting surface sites can be subsequently modified chemically to provide surface charges. Examples of such species include N-2-aminoethyl-3-aminopropyltrimethoxy-silane, 3-aminopropyltrimethoxysilane, 3-methacryloxypropyl-trimethoxysilane, and vinyltriacetoxysilane. Alternatively, other species can be simply adsorbed to the surface of the inorganic material particles. The most important examples of these are low molecular weight polyelectrolytes such as conventional pigment dispersants.

Examples of suitable anionic polyelectrolyte pigment dispersants for use in the process of the present invention include polyacrylic acid, polymethacrylic acid, copolymeric acids including copolymerized maleic acid, polyphosphates such as potassium tripolyphosphate, and the like.

While the chemical composition of a polymeric latex binder is important for achieving the resultant properties

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of the coating or film when the polymeric latex also acts as the binder, the glass transition temperature and amount of acid functionality in the polymeric latex can also be important for the purpose of the invention. Hard, irreversibly absorbing polymeric latex particles, preferably having an average particle diameter of from about 50 nm to 80 nm, are preferred in one embodiment of this invention. Soft, irreversably adsorbing polymeric latex particles preferably having a particle diameter of from about 100 nm to about 400 nm are preferred in another embodiment. 10 Further, the amount of acid functionality at the surface of the polymeric latex particles is believed to be especially important. In a presently preferred embodiment of the present process, a polymeric latex is employed by a process which preferentially provides acid functionality at the 15 surface of the polymer latex particles. Examples of polymerization processes which preferentially provide surface acid are described in K. L. Hoy, J. Coat. Tech. 51 27-41 (1979).

20 The presence of conventional polyelectrolyte dispersants has a significant effect on the adsorption of the selected polymeric latex on the inorganic material particles in the process of the present invention. low concentrations of conventional water-soluble polyelectrolyte dispersants, for example in the case of 25 titanium dioxide, on the order of about 0.2 weight percent or lower, have little adverse effect on the adsorption of selected polymeric latexes onto inorganic material particles surfaces, the use of higher concentrations of these conventional water-soluble polyelectrolytes dispersants can 30 have a significant adverse effect upon adsorption unless the polymeric latex is carefully selected.

In some cases, it is possible to practice the process of the present invention without employing a pigment dispersant to disperse the inorganic material particles, although it is generally preferred that a pigment dispersant

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be used.

In one presently preferred embodiment of the present invention, an emulsion polymerization process known to provide acid functionality preferentially at the surface of the polymeric latex particles is employed to prepare the 5 polymeric latex for adsorption on the inorganic material particles, such as titanium dioxide particles. case, the level of polyelectrolyte dispersant in the pigment dispersion and the formulated coating can be higher, such as up to one percent or more of the weight of the inorganic 10 material depending on the specific polyelectrolyte dispersant employed. In this case, a weakly adsorbing polyelectrolyte dispersant, such as Tamol® (trademark of Rohm and Haas Company) SG-1 dispersant (ammonium salt of copolymer of a carboxylic acid monomer and a hydrophilic 15 comonomer) can be employed at a high level such as one percent, while a strongly adsorbing polyelectrolyte dispersant such as a polyacrylic acid dispersant, such as Tamol 731 dispersant (sodium salt of polymeric carboxylic acid), which strongly adsorbs to the surface of the titanium dioxide particles, and which displaces weakly adsorbed polyelectrolyte dispersants from the surface of titanium dioxide particles, is preferably used at low levels.

In another presently preferred embodiment of the present invention, an emulsion polymerization process is 25 used to prepare a polymeric latex from monomer mixture including one or more ethylenically polymerizable derivatives of phosphoric acid, such as one or more partial esters of phosphoric acid and 2-hydroxyethyl methacrylate. Examples of such partial esters are given above and include 30 the monoester of phosphoric acid with hydroxyethyl methacrylate, the diester of phosphoric acid with hydroxyethyl methacrylate, and mixture thereof. The resulting phosphoric acid ester-functional polymeric latex particles adsorb to inorganic material particles more 35 strongly than polymeric latex prepared from monomer mixture

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which does not include phosphoric acid ester-functional monomer.

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The process of the present invention can significantly increase the hiding of titanium dioxide containing emulsion paint formulations (as determined by measuring the scattering coefficient of the paint) of the same PVC compared with the scattering coefficient of a paint formulation prepared by the use of conventional polyelectrolyte dispersants. An alternate way of expressing this improvement is that the process of the invention permits the reduction of the PVC of a 20% PVC titanium dioxide paint formulation by 20 percent while achieving the same scattering coefficient or degree of hiding as can be obtained by the use of a conventional polyelectrolyte pigment dispersant.

Furthermore, the improved performance of paint formulations prepared by the present process is achieved even in the presence of substantial amounts of large particle size extenders, such as calcium carbonate and alumino silicate, typically employed in such formulations. In this regard the improved hiding is obtained by the controlled adsorption of the primary pigment, the titanium dioxide. This improvement is not lost upon the incorporation of other types of inorganic or organic particles, such as extender pigment particles.

In addition, the improved hiding achievable by use of the process of this invention is realized when the coating formulation is prepared with conventional thickeners of the associative and non-associative types; this improvement is not dependent upon the selected thickener. This result is surprising and contrary to the result typically encountered when paints are formulated using conventional polyelectrolyte dispersants. In such conventional systems the hiding property of the formulated paint can vary greatly as a function of the thickener employed. The controlled adsorption process of the invention therefore offers paint

formulators a greater choice in selecting thickeners for the final paint formulation without concern over the choice of thickener adversely affecting the final hiding properties of the paint.

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Furthermore, in addition to improving the hiding of paint formulations, the process of this invention also has been found to advantageously improve the early blister resistance, metal marking resistance, gloss, high shear viscosity, and scrub resistance of paint formulations compared with paint formulations dispersed with conventional polyelectrolyte dispersants, as well as improving the corrosion resistance of paint formulations.

In addition to the titanium dioxide particles, which are typically of relatively small particle size, on the order of 200-250 nm, other pigment particles, such as the large particle size extender pigment particle typically included in many types of coating formulation to reduce new material costs, can also be employed in addition to the titanium dioxide pigment particles in the process of the present invention. Examples of large particle size extender pigment particles which can be used include calcium carbonate, alumio-silicate, amorphous silicon, and the like.

Similarly, the aqueous medium in which the titanium dioxide particles are ground with the polymeric latex dispersant can also include water-miscible solvents, such as glycols and glycol ethers, such as are conventional in the coatings arts. Examples of water-miscible solvents employed include propylene glycol, ethylene glycol, ethylene glycol monomethyl ether, and the like.

Aqueous coating compositions are frequently formulated at alkaline pH to stabilize anionically charged latex binder against agglomeration and for other reasons. The principles of formulating aqueous coating compositions are reviewed, for example, in <u>Formulation of Organic Coatings</u> (N.I. Gaynes ed. D. Van Nostrad Co. Inc. Princeton, NJ 1967) at pp.

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The present invention provides aqueous dispersions of composite particles which can be used in a variety of applications. In some cases, a "soft" (i.e. low effective glass transition temperature, such as on the order of 0 °C to 10 °C) polymeric latex can be added to the dispersion of composite particles, typically with the further addition of other components, the soft polymeric latex serving as binder for the composition. Often the binder forming latex polymer particles contain carboxyl functional groups. Under suitable conditions, the carboxyl groups are ionized and the resultant charges on the latex particle surface electrostatically stabilize the latex against premature agglomeration. Often a volatile base, typically ammonia, is used to adjust the pH of the coating composition. When the coating composition is applied to a substrate to be coated, the volatile base is lost and the pH of the coating composition drops, destabilizing the latex particles of the binder and thus encouraging agglomeration to form a continuous binder film.

In addition to binder-forming polymeric latex particles and composite polymeric latex-inorganic material particles, aqueous coating compositions prepared according to the process of the present invention can include typical coatings ingredients. For example, they can include extender pigments as noted above such as calcium carbonate, amorphous silica, and the like; defoamers; biocidal agents such as zinc oxide, 2-N-octyl-4-isothiazole-3-one, and phenyl mercuric acetate; coalescing agents such as diethylene glycol monoethyl ether acetate and ethylene glycol monobutyl ether acetate; plasticizers such as dialkyl phthalates including dibutyl phthalate, dioctyl phthalate and the like; freeze-thaw stabilizers such as ethylene glycol, propylene glycol and diethylene glycol; nonionic wetting agents such as polyethylenoxylates of fatty acids, alkanols, alkylphenols, and glycols; polyelectrolyte pigment

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dispersants at low levels; thickeners such as polyelectrolyte and cellulosic thickeners; rheology control agents such as associative thickeners and the like; colorants such as colored pigments and dyes; perfumes; cosolvents and the like.

In the examples which follow, the following abbreviations are used:

	AA	acrylic acid
	MAA	methacrylic acid
10	IA	itaconic acid
	EA	ethyl acrylate
	BA	butyl acrylate
	MMA	methyl methacrylate
	PEM	phosphoethyl methacrylate

The illustrative examples which follow illustrate the process of the present invention as a function of the parameters of the selected emulsion polymer and the adsorption achieved. These examples will aid those skilled in the art in understanding the present invention; however, the present invention is in no way limited thereby. In the examples which follow, percentage composition is by weight, unless otherwise noted.

Examples 1 - 3

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A typical conventionally dispersed 20% PVC gloss paint (Comparative Example 1) was prepared according to the following formulation. Pigment grind:

30	Components	Weight (g)
	water	30
	propylene glycol	20
	Colloid 643 defoamer	1
	Tamol® 731 dispersant (25% solids)	8.19
35	Ti-Pure® R-900 titanium dioxide	204.72

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Tamol 731 is a polycarboxylate dispersant (Tamol is a trademark of Rohm and Haas Company). Tamol 731 is a sodium salt of polymeric carboxylic acid. Pigment dispersion with one percent by weight polymeric carboxylic acid dispersant as in this case is considered conventional. Ti-Pure is a trademark of Du Pont de Nemours Co. Ti-Pure R-900 is a coatings grade of rutile titanium dioxide. These components were milled on a high speed disk disperser to form a pigment grind, and were let down at a slower speed with the following:

	Components	Weight (g)
	Rhoplex® AC-61 polymer latex	493.75
	Colloid 643 defoamer	4
15	Texanol® coalescent	22.96
	Triton® GR-7M surfactant	2
	propylene glycol	59
	Nuosept® 95 preservative	6
	water and Natrosol® 250 MR	165.07
20	cellulosic thickener	

Rhoplex is a trademark of Rohm and Haas Company. Texanol is a trademark of Eastman Kodak Co. Triton is a trademark of Rohm and Haas Company. Nuosept is a trademark of Nuodex, Inc. The mix is thickened to a Stormer viscosity of 80 KU by addition of an aqueous solution of the cellulosic thickener, and the pH of the mix is adjusted to 9 by addition of ammonia.

To provide Example 1, the paint formulation process of Comparative Example 1 was repeated replacing the 493.75 g of Rhoplex AC-61 with a mixture of 419.66 g Rhoplex AC-61 (46.5% solids) and 92.04 g of a polymeric latex having a glass transition temperature of 65°C, a particle size of 71 nm, and having 6% phosphoethyl methacrylate (Latex A, 37.4% solids). The total weight of water was adjusted to maintain the same total solids in the final paint.

The paint formulation process of Example 1 was

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repeated with the 1% Tamol 731 decreased to 0.5% to give Example 2.

The paint formulation process of Example 1 was repeated substituting Tamol SG-1 for Tamol 731 to give Example 3.

For each paint, a sample was removed just before addition of the cellulosic thickener, allowed to equilibrate for two hours, diluted 1 part to 3 parts with water, and centrifuged. The concentration of unadsorbed latex in the supernatant was determined gravimetrically and used to calculate the milligrams of latex adsorbed per gram of titanium dioxide.

Kubelka-Munk scattering coefficients were determined for all paints using a modification of ASTM D 2805-70.

Results given in Table I.

Table I

				Adsorption	Scattering
	Example	<u>Dispersant</u>	<u>Binder</u>	(mg/g)	<u>Coefficient</u>
20	Comp. 1 ¹	1% Tamol 731	AC-61	3	7.26
	1		AC-61/Latex	A 199	7.77
	2	0.5% Tamol 731	AC-61/Latex	A 200	8.18
	3	1% Tamol SG-1	AC-61/Latex	A 225	8.27

²⁵ Comparative example.

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The results for Example 1 in Table I demonstrate that a polymeric latex with high acid functionality at the latex particle surface when blended with a conventional latex will preferentially adsorb on titanium dioxide under conditions where the conventional latex will not adsorb. Thus Comparative Example 1 shows that Rhoplex AC-61 does not adsorb on titanium dioxide in the presence of 1% Tamol 731 while Example 1 shows that a blend of Latex A with Rhoplex AC-61 provides substantial adsorption and increased hiding as reflected by the Kubelka-Munk scattering coefficients.

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The latex adsorption of 199 mg/g measured for Example 1 is close to value of 169 mg/g measured independently for Latex A in the absence of Rhoplex AC-61 and is significantly below the value of about 400 mg/g measured independently for Rhoplex AC-61. The lower adsorption observed for Latex A is consistent with its smaller 71 nm particle size compared to the 150 mm Rhoplex AC-61.

Comparison of the results for Examples 2 and 3 with those for Example 1 shows that amount and type of dispersant used to prepare the titanium dioxide pre-disposed has a significant effect on the behavior of the paint made by letting down this pre-dispersion with a blend of Rhoplex AC-61 and polymeric Latex A. Either decreasing the amount of Tamol 731 (Example 2) or replacing the Tamol 731 with Tamol SG-1, which has a lower proportion of acid functionality and is believed to be more weakly bound to the titanium dioxide surface (Example 3), provides greater adsorption and increased hiding.

Examples 4 - 9

A series of polymeric latexes of varying particle size and glass transition temperature and containing varying amounts of acidic monomer, as given in Table II, were prepared. Next, a series of 35% PVC titanium dioxide grinds were prepared using Ti-Pure R-900 titanium dioxide dispersed with 0.05% by weight Tamol 731 dispersant. In preparing these grinds, a conventional Tamol 731-dispersed grind was prepared first; polymeric latex was then added and the mixture milled on a high speed disk disperser for 15 minutes.

The polymeric latexes were prepared using a gradual-addition, batch process, with all of the acidic monomer being added in the monomer mixture, except in the case of Example 6, in which the acidic monomer was added in a manner known to provide a preferred distribution of acid

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functionality at the surface of the polymeric latex.

Immediately after grinding a sample was removed, diluted 1 part to 3 parts with water, and centrifuged. The concentration of unadsorbed latex in the supernatant was determined gravimetrically and used to calculate the milligrams of latex adsorbed per gram of titanium dioxide.

To determine the reversibility of the latex adsorption, 1% polyacrylic acid (Tamol 963 dispersant) was added to a portion of the remaining grind and the mixture stirred for 15 minutes. After equilibrating for 1 week, latex adsorption was determined as before. This procedure was carried out immediately after the grind was made and after it had been allowed to equilibrate for 1 day and 1 week. The results are given in Table II.

These results in Table II show that the ability of polyacrylic acid to desorb previously adsorbed latex decreases with increasing surface acidity of the latex.

Surface acidity is controlled in these Examples 4-9 by varying the level of acidic monomer (Comparative Example 3 and Examples 4-5), or by varying the polymerization process to enhance the surface acid (Example 6), but other methods of influencing latex surface acidity should be equally satisfactory.

The results in Table II also demonstrate that the type
of acid functionality greatly influences reversibility.
Thus replacing methacrylic acid with phosphoethyl
methacrylate results in a decrease in desorption (Examples 4
and 8). A similar decrease in reversibility is obtained
with itaconic acid (Example 9).

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Table II

	Ex.	Part. Size (nm)			Total Latex (mg/g)	Adsorption (mg/g)		Equili	ption ³ brat. <u>1Week</u>
5	4	64	51	5% MAA	158	137	91	67	20
	5	66	52	10% MAA	158	132	95	7	2
	6	64	51	5% MAA	158	137	64	1	0
	7	73	94	5% MAA	158	152	109	105	97
	8	56	54	2% MAA-	+ 158	156	23	7	3
10				2% PEM ¹					
	9	69	50	2% IA ²	158	157	22	- 1	- 1.

^{1.} PEM = phosphoethyl methacrylate.

20 Example 10

A typical conventionally dispersed pigment grind containing one percent Tamol SG-1 polymeric carboxylic acid dispersant by weight on pigment was prepared according to the following formulation:

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	Components	Weight	<u>(g)</u>
	water	70	
	propylene glycol	20	
	Colloid 643 defoamer	1	
30	Tamol SG-1 dispersant (35% solids)	5.85	
	Ti-Pure R-900 titanium dioxide	204.72	

These components were milled on a high speed disk disperser to form a pigment grind. 237.6 g of a 120 nm polymeric latex (38.0% solids) prepared using a gradual-addition, batch process and having a monomer composition of 5 BA/93 MMA/2 MAA was then added and the

^{15 2.} IA = itaconic acid.

^{3.} Desorption values less than 0 and greater than 100 reflect experimental error in the desorption measurement and imply no desorption and complete desorption respectively.

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mixture milled for 15 minutes to give Comparative Example 2.

After equilibrating for 1 hour, a sample was removed, diluted 1 part to 3 parts with water, and centrifuged. The concentration of unadsorbed latex in the supernatant was determined ravimetrically and used to calculate the milligrams of latex adsorbed per gram of titanium dioxide. The resulting value for Comparative Example 2 was 25 mg/g.

Example 10 was prepared in the same manner using 239.5 g of a 127 nm polymeric latex (37.7% solids) prepared using a gradual addition, batch process and having a monomer composition of 5 BA/89 MMA/2 MAA/4 phosphoethyl methacrylate. The measured adsorption for Example 10 was 219 mg/g.

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A portion of each grind was diluted to about one percent by volume with water, a small drop placed on a standard SEM sample holder, and the water allowed to evaporate. The dried samples were then sputter coated with gold and examined via scanning electron microscopy. The resulting micrographs (magnification of 30,000:1) are shown in Figure 4a (Example 10) and Figure 4b (Comparative Example 2).

Figure 4a shows "raspberry"-shaped composite particles resulting from the adsorption of the small latex particles on the surfaces of the larger titanium dioxide particles. By contrast, the smooth titanium dioxide particles and the unadsorbed latex particles seen in Figure 4b demonstrate the absence of such adsorption for a conventional phosphoethyl methacrylate-free latex. These micrographs provide a direct visual confirmation of the adsorption process of the present invention.

Example 11

An acrylate polymeric latex emulsion was prepared having a calculated glass transition temperature of about 14 °C, a particle size of 161 nm, and total solids of about 45

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percent by weight using a single-stage, gradual-addition thermal process from monomer including 1.3 percent by weight methacrylic acid and one percent by weight phosphoethyl methacrylate (Latex B). A pigment grind was prepared by grinding together at high speed 200 parts by weight TiPure R-900 titanium dioxide, 8 parts by weight Tamol 731 pigment dispersant, 1 part by weight Colloid 643 defoamer, 20 parts by weight propylene glycol and 30 parts by weight water. 495.6 parts by weight of Latex B were mixed at low speed with 7.57 parts by weight water and 22.64 parts by 10 weight Texanol. Subsequently, to the Latex B/Texanol mixture were added 9.70 parts by weight Triton X-405 surfactant, 49.57 parts by weight water, 4 parts by weight Colloid 643 defoamer, 59 parts by weight propylene glycol, and 6 parts by weight Nuosept 95 preservative. To this 15 mixture was added the grind, and subsequently, 100.4 parts by weight of a 5.0 percent by weight solution of Natrosol 250 MR hydroxycellulose thickener were added to provide a paint (Example 11) with a PVC of 20, a calculated volume solids of 32 percent, and a calculated weight solids of 42.1 20 percent. The hiding and gloss of the paint were evaluated and compared with a control paint prepared from an polymeric latex made using a similar process but omitting the phosphoethyl methacrylate (Comparative Example 3), the results being given in Table III, and showing improved 25 hiding and gloss using the process of the present invention.

Table III

30	<u>Example</u>	<u>Hiding</u>	20° Gloss	60° Gloss
-	Comp. 31	6.9	14	54
	11	7.8	24	63

^{1.} Comparative example.

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Examples 12 - 13

A multi-stage gradual addition process was employed to prepare an acrylate copolymer emulsion from monomer including 1.9 percent by weight methacrylic acid. The 5 polymeric latex, Latex C, contained 44.7% solids with an average particle diameter of 166 nm. Latex C is believed to have a substantial proportion of acid functionality at the surface of the latex particle as determined by potentiometric titration. A modification of the process was 10 employed to prepare an acrylate copolymer emulsion, Latex D, from monomer having the same composition but which is believed to result in an emulsion polymer having a substantially lower proportion of acid functionality at the polymer surface. Latex D contained 44.6% solids with an 15 average particle diameter of 158 nm.

A tinted titanium dioxide pre-dispersion in which the dispersant (Tamol SG-1 dispersant) level was 1% (on titanium dioxide) was prepared according to the following formulation.

	Component	Weight (g)
	water	169.9
	Nopco NXZ defoamer	3.0
25	Tamol SG-1(35%) dispersant	17.1
	Ti-Pure R-900 titanium dioxide	600

The pigment grind components were milled on a high speed disk disperser (Laboratory Dispersator, Model 84, Premier Mill Corp.) at 3000 rpm for 20 minutes. At a lower speed 30.0 g of Colortrend (trademark of Tenneco Inc.) 888-9907 B-Lamp Black was mixed in.

To 62.9 g of the tinted pre-dispersion was added with stirring 169.4 g of Latex C binder (preadjusted to pH 6.0 with 28% aqueous ammonia) and 9.3 g of water. To 120.8 g of the resulting mixture was added 5.2 g Texanol coalescent,

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7.2 g propylene glycol, and 24.0 g of a 2.5% aqueous solution of Natrosol 250 MR. The final pH of this mixture was adjusted to 7.5 with 28% aqueous ammonia to yield a paint (Example 12) with 14% volume concentration of titanium dioxide in the dry paint and a volume solids content of ca. 30% in the wet paint.

The above procedure was repeated using 169.7 g of Latex D binder (preadjusted to pH 6.0) instead of Latex C and 9.0 g of water to form an analogous paint (Comparative Example 4).

A tinted pigment dispersion was prepared as in Example 12 except that 17.3 g of Tamol 963 dispersant (35%) (sodium salt of polyacrylic acid) instead of Tamol SG-1 dispersant was used to give a one percent Tamol 963 dispersion.

The paint formulation process of Example 12 was repeated substituting the Tamol 963 pigment dispersion for the Tamol SG-1 pigment dispersion using Latex C to provide Example 13 and Latex D to provide Comparative Example 5.

After four days the paints were evaluated for latex adsorption and for differences in titanium dioxide scattering efficiency. Adsorption was measured by diluting one part of the paint with seven parts by weight of water, centrifuging, and determining the concentration of the unadsorbed latex in the supernatant gravimetrically.

The relative scattering was determined by measuring the Y-reflectance of the tinted paints with a 45'/0' reflectometer (Colorguard, Gardner Instruments). From light scattering theory, the ratio of the scattering coefficient, S, and the adsorption coefficient, K, is given by:

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$$S/K = 2R/(1-R)^2$$

where R is the reflectance from an infinitely thick layer of dry paint film. At a given titanium dioxide PVC,

35 differences in Y-reflectance of equally tinted paint films arise solely from differences in titanium dioxide scattering

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efficiency. Titanium dioxide dominates the scattering while the tinting agent dominates the adsorption which can then be considered constant. The percentage difference in scattering coefficient exhibited by two equally tinted paints is then given by:

$$((S_2 - S_1)/S_1) \times 100 = ((R_2(1 - R_1)^2/R_1(1 - R_2)^2) - 1) \times 100$$

Results are given in Table IV. The results in Table IV show that the Latex C polymeric binder (Example 12) provides 10 substantially higher adsorption on titanium dioxide than does the Latex D polymeric binder (Comparative Example 4). The difference in Y-reflectance represents 48% higher scatter from the titanium dioxide. This result demonstrates the advantage of a high acid functionality at the latex 15 particle surface on the adsorption process of this invention. High surface acid functionality is believed to promote adsorption to titanium dioxide thereby enhancing the ability of the latex binder to compete with the polyelectrolyte dispersant for the titanium dioxide surface. The results for Example 13 and Comparative Example 5 show no difference between the two in adsorption and essentially no difference in Y-reflectance.

Comparison of the results for Examples 12 and 13 show that the dispersant employed to prepare the titanium dioxide predispersion has a significant effect in this case on the adsorption of the polymeric latex dispersant Latex C with high surface acid functionality. When the polyelectrolyte Tamol 963 dispersant was employed, it was not displaced from the surface of the titanium dioxide particles to permit adsorption of the polymeric Latex C. However, this same polymeric latex was effective to displace the Tamol SG-1 dispersant, which has a lower proportion of acid functionality on a weight basis than the Tamol 963 dispersant and is believed to be more weakly bound to the titanium dioxide surface.

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Table IV

				Adsorption	
	<u>Example</u>	<u>Latex</u>	Dispersant	(mg/g)	Y-Reflectance
5	12	С	1% Tamol SG-1	720	0.557
	$C. 4^{1}$	D	1% Tamol SG-1	110	0.492
	13	С	1% Tamol 963	50	0.544
	C. 5 ¹	D	1% Tamol 963	50	0.541

10 1. Comparative example.

Example 14

A multi-stage gradual addition process was employed to 15 prepare an acrylate copolymer emulsion from monomer including about 2.5 percent by weight methacrylic acid. The polymeric latex, Latex E, contained 44.5% solids with an average particle diameter of 150 nm, and was prepared by a process believed to provide a substantial proportion of acid . functionality at the surface of the latex particle as 20 determined by potentiometric titration. A modification of the process was employed to prepare an acrylate copoymer emulsion, Latex F, from monomer having the same composition, but which is believed to result in an emulsion polymer having a substantially lower proportion of acid 25 functionality at the polymer surface. The latex, Latex F, contained 44.8% solids with an average particle diameter of 159 nm.

To 86.1 of Latex E (preadjusted to pH 9.0 with aqueous ammonia) was added 31.5 g of a titanium dioxide-Tamol SG-1 dispersion prepared as in Examples 12-13, 3.3 g of deionized water, 1.9 g of Texanol coalescent, 7.2 g of propylene glycol, and 24.0 g of a 2.5% aqueous solution of Natrosol 250 MR. These ingredients were well mixed to yield a paint (Example 14) with 14% PVC of titanium dioxide and volume solids of ca. 30%.

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To 85.2 g of Latex F (preadjusted to pH 9.0 with aqueous ammonia) was added 31.5 g of the same titanium dioxide-Tamol SG-1 dispersion, 6.2 g of deionized water, 1.9 g of Texanol coalescent, 7.2 g of propylene glycol, and 24.0 g of a 2.5% aqueous solution of Natrosol 250 MR. These ingredients were well mixed to yield a paint (Comparative Example 8) with 14% PVC of titanium dioxide and volume solids of ca. 30%.

Example 14 and Comparative Example 6 were evaluated for polymeric latex adsorption after one hour, one day, and five days of equilibration. Both paints were also evaluated for Y-Reflectance after one day and nine days of equilibration in the same manner as Examples 12 - 13. The results are given in Table V.

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Table V

			Adsorpt	ion (g/	g TiO_2)	Y-Refl	ectance
	<u>Example</u>	<u>Latex</u>	1 hour	1 day	5 days	1 day	9 days
20	14	E	0.01	0.17	0.23	0.530	0.536
	Comp. 61	F	-0.05	-0.09	-0.03	0.502	0.498

^{1.} Comparative example.

25 These results demonstrate that in case of a polymeric latex with high acid functionality at the latex particle surface, such as Latex E, the adsorption process is not instantaneous but occurs over the time scale of days. After one day a paint prepared with a high surface acid functionality polymeric latex binder, Latex E, had 18.5% higher light scatter than a paint prepared with a low surface acid functionality polymeric latex, Latex F. After

nine days this difference increased to 26%.

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Example 15

To 1300 g of deionized water stirred under a nitrogen atmosphere at 85°C was added 4 g of ammonium persulfate in 24 g of deionized water and 60 g of a seed latex. A monomer emulsion prepared from 600 g of deionized water, 28 g of Siponate DS-4, 900 g of butyl acrylate, 876.6 g of methyl methacrylate and 23.4 g of methacrylic acid was added over a 200-min period along with 2 g of ammonium persulfate in 100 g of deionized water maintaining 85°C. When the reaction was complete, the dispersion was cooled and filtered. The product (Latex G) had 45.6% solids and pH 3 with an average diameter of 196 nm.

The process used to prepare Latex G was repeated, except that 18 g of the methyl methacrylate was replaced with an equal weight of Kayamer® (trademark of Nihon Kayaku Co. Ltd.) PM-1 (monomer mixture containing around 52% of the monoester of hydroxyethyl methacrylate and phosphoric acid and 33% of the diester). The product (Latex H) had 46.1% solids and pH 2.6 with an average particle size of 185 nm.

A tinted titanium dioxide dispersion was prepared as in Examples 12-13 from 180 g of deionized water, 7.2 g of Tamol 731 dispersant (25% solids), 3.0 g of Nopco NXZ defoamer, 600 g of Ti-Pure R-902 and 30 g of Colortrend lamp black dispersion. Samples of Latex G and Latex H were adjusted to pH 9 with 28% aqueous ammonia and used to formulate paints with 14% volume concentration of Ti-Pure R-902 pigment in the dry paint: To 29.5 g of R-902 dispersion was added 7.8 g of water, 83.9 g of the pH 9 Latex G, 2.6 g of Texanol coalescent, 7.2 g of propylene glycol and 24.0 g of 2.5% aqueous Natrosol 250 MR hydroxyethyl cellulose thickener: Latex H was formulated similarly only 82.8 g were required and 8.9 g of water. The two paints were allowed to equilibrate for seven days and then the adsorption of polymeric latex on titanium dioxide in the paints and their Y-reflectance was measured as above.

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The results are given in Table VI.

Table VI

5	_Example_	<u>Latex</u>	Adsorption (g/g TiO2)	Y-Reflectance
	Comp. 71	G	0.01	0.504
	15	H	0.26	0.540

1. Comparative example

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These results show that the standard binder (Latex G) is essentially non-adsorbing under these conditions. The phosphate-containing binder (Latex H) adsorbs substantially and the increased Y-reflectance that accompanies the adsorption corresponds to an increase in scattering efficiency of the titanium dioxide in the dry paint film of 25%.

Various modifications can be made in the details of the various embodiments of the compositions and processes of the present invention, all within the spirit and scope of the invention as defined by the appended claims.

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CLAIMS

- 1. A process for preparing an aqueous dispersion of composite particles, the composite particles each comprising a plurality of selected polymeric latex particles adsorbed onto an inorganic material particle, the process comprising:
- a) dispersing inorganic material particles in an aqueous medium;
- b) preparing a dispersion or suspension of selected polymeric latex particles in the aqueous medium, the polymeric latex particles being prepared by an emulsion polymerization process, the emulsion polymerization process being selected from the class consisting of:
- 1) emulsion polymerization processes employing a monomer mixture including at least one polymerizable
 15 ethylenically unsaturated acid-functional monomer, the at least one polymerizable ethylenically unsaturated acid-functional monomer being selected from the class consisting of:
 - A) itaconic acid; and
- B) dihydrogen phosphate esters of an alcohol, the alcohol containing a polymerizable olefinic group;
 - 2) emulsion polymerization processes preferentially providing acid-functionality proximate the surface of the polymeric latex particle; and
- 3) emulsion polymerization processes employing a monomer mixture including at least one ethylenically unsaturated acid-functional monomer, the total weight of ethylenically unsaturated acid-functional monomer comprising at least about ten percent by weight of the polymeric latex solids; and
 - c) mixing the aqueous medium containing the inorganic material particles and the selected polymeric latex particles, the selected polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles.
 - 2. A process according to claim 1 wherein the

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medium by employin pigment dispersant, the pigment dispersant adsorbin not the inorganic material particles, and the selected possible in the presence of the pigment dispersant to provide the composite particles when the aqueous medium containing the inorganic material particles and the selected polymeric latex particles is mixed.

- 3. A process according to claim 2 wherein the dihydrogen phosphate ester is selected from the phosphoric acid monoester of 2-hydroxyethyl methacrylate, the phosphoric acid diester of 2-hydroxyethyl methacrylate, the phosphoric acid monoester of 3-hydroxypropyl methacrylate, and the phosphoric acid diester of 3-hydroxypropyl methacrylate.
 - 4. A process according to claim 2 wherein the pigment dispersant is a polyelectrolyte.
- 5. A process according to claim 4 wherein the pigment dispersant is selected from poly(meth)acrylic acid, poly(meth)acrylic acid salts, polyelectrolyte copolymers of (meth)acrylic acid and salts of polyelectrolyte copolymers of (meth)acrylic acid.
- 6. A process according to claim 4 wherein the pigment dispersant is an inorganic polyelectrolyte.

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- 7. A process according to claim 6 wherein the pigment dispersant is a polyphosphate salt.
- 8. A process according to claim 2 wherein the selected polymeric latex has an effective glass transition temperature greater than about 20 °C.
- 9. A process according to claim 8 wherein the selected polymeric latex has an effective glass transition temperature greater than about 50 °C.
- 10. A process according to claim 2 wherein the average 35 size of the polymeric latex particles is from about 20 nm to about four times the average particle size of the inorganic

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material particles.

- 11. A process according to claim 10 wherein the average size of the polymeric latex particles is from about one-half the average size of the inorganic material particles to about equal to the average size of the inorganic material particles.
- 12. A process according to claim 2 wherein the average particle size of the inorganic material particles is from about 100 nm to 10,000 nm, and the average size of the polymeric latex particles is from about 20 nm to 5000 nm.
- 13. A process according to claim 12 wherein the average particle size of the inorganic material particles is from about 200 nm to 250 nm, and the average size of the polymeric latex particles is from about 20 nm to 1000 nm.
- 14. A process according to claim 2 wherein the effective glass transition temperature of the selected polymeric latex is less than about 20 °C, the average particle size of the selected polymeric latex being from about 80 nm to 600 nm.
- 15. A process according to claim 14 wherein the average particle size of the selected polymeric latex is from about 100 nm to 400 nm.
 - 16. A process according to claim 8 wherein the average particle size of the polymeric latex particles is from about 30 nm and 100 nm.
 - 17. A process according to claim 16 wherein the average particle size is from about 50 nm to 80 nm.
 - 18. A process according to claim 2 wherein the inorganic material is titanium dioxide.
- 19. A coating composition produced according to the process of claim 2.
 - 20. A process for preparing an aqueous composition, the composition including dispersed composite particles, the composite particles each comprising a plurality of selected polymeric latex dispersant particles adsorbed to a inorganic material particle, the process comprising:

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- a) dispersing inorganic material particles in an aqueous medium employing a pigment dispersant, the first pigment dispersant adsorbing onto the inorganic material particles;
- b) preparing a dispersion or suspension of selected polymeric latex particles in the aqueous medium, the polymeric latex particles being prepared by an emulsion polymerization process, the emulsion polymerization process being selected from the class consisting of:
- 1) emulsion polymerization processes employing a monomer mixture including at least one polymerizable ethylenically unsaturated acid-functional monomer, the at least one polymerizable ethylenically unsaturated acid-functional monomer being selected from the class consisting of:
 - A) itaconic acid; and
 - B) dihydrogen phosphate esters of an alcohol, the alcohol containing a polymerizable olefinic group;
 - 2) emulsion polymerization processes preferentially providing acid-functionality proximate the surface of the polymeric latex particle; and
 - 3) emulsion polymerization processes employing a monomer mixture including at least one ethylenically unsaturated acid-functional monomer, the total weight of ethylenically unsaturated acid-functional monomer comprising at least about ten percent by weight of the polymeric latex solids; and
 - c) mixing the aqueous medium containing the inorganic material particles and the selected polymeric latex particles, the selected polymeric latex particles adsorbing onto the inorganic material particles in the presence of the pigment dispersant to provide the composite particles.
- 21. A process according to claim 20 wherein the dihydrogen phosphate ester is selected from the phosphoric acid monoester of 2-hydroxyethyl methacrylate, the

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phosphoric acid dies of 2-hydroxyethyl methacrylate, the phosphoric acid mono of 3-hydroxypropyl methacrylate, and the phosphoric accordance of 3-hydroxypropyl methacrylate.

- 22. A process ac ling to claim 21 further comprising mixing an aqueous dispersion of polymeric latex binder particles with the aqueous medium containing the dispersed composite particles.
- 23. An aqueous dispersion of composite particles, the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including dihydrogen phosphate ester functional groups.
 - 24. An aqueous dispersion according to claim 23, the polymeric latex particles being polymerized from monomer including at least one polymerizable dihydrogen phosphate-functional monomer.
- 25. An aqueous dispersion according to claim 24, the at least one polymerizable dihydrogen phosphate-functional monomer being selected from the dihydrogen phosphate esters of an alcohol, the alcohol including a polymerizable group selected from polymerizable vinyl groups and polymerizable non-vinyl olefinic groups.
- 26. An aqueous dispersion according to claim 25 wherein the at least one polymerizable dihydrogen phosphate-functional monomer is selected from the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate, and mixtures of the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate and the phosphoric acid diester of 2-hydroxyethyl methacrylate.
 - 27. An aqueous dispersion according to claim 25 wherein the at least one polymerizable dihydrogen phosphate-functional monomer is selected from the dihydrogen vinylbenzylphosphates, and CH₂=CCH₃CO₂CH₂CH₂OPO (OH)₂,
- CH₂=CCH₃CO₂CH₂CH [OPO (OH)₂] CH₃, CH₂=CCH₃CO₂CH₂CH₂CH₂OPO (OH)₂, and CH₂=CCH₃CO₂CH₂CHOHCH₂OPO (OH)₂.

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- 28. A coating composition comprising an aqueous dispersion of composite particles, the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including dihydrogen phosphate ester functional groups.
- 29. A coating composition according to claim 28, the polymeric latex particles being polymerized from monomer including at least one polymerizable dihydrogen phosphate-functional monomer.
- 30. A coating composition according to claim 30, the at least one polymerizable dihydrogen phosphate-functional monomer being selected from the dihydrogen phosphate esters of an alcohol, the alcohol including a polymerizable group selected from polymerizable vinyl groups and polymerizable non-vinyl olefinic groups.
- 31. A coating composition according to claim 29 wherein the at least one polymerizable dihydrogen phosphate-functional monomer is selected from the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate, and mixtures of the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate and the phosphoric acid diester of 2-hydroxyethyl methacrylate.
- 32. A coating composition according to claim 30 wherein the at least one polymerizable dihydrogen phosphate-functional monomer is selected from the dihydrogen vinylbenzylphosphates, and CH₂=CCH₃CO₂CH₂CH₂OPO (OH)₂, CH₂=CCH₃CO₂CH₂CH₂CH₂OPO (OH)₂, and CH₂=CCH₃CO₂CH₂CH₂CH₂CHOHCH₂OPO (OH)₂.
- 33. A pigment slurry comprising an aqueous dispersion of composite particles, the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including dihydrogen phosphate ester functional groups.
 - 34. A pigment slurry according to claim 33, the

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polymeric latex particles being polymerized from monomer including at least one polymerizable dihydrogen phosphate-functional monomer.

- 35. A pigment slurry according to claim 34, the at least one polymerizable dihydrogen phosphate-functional monomer being selected from the dihydrogen phosphate esters of an alcohol, the alcohol including a polymerizable group selected from polymerizable vinyl groups and polymerizable non-vinyl olefinic groups.
- 36. A pigment slurry according to claim 35 wherein the at least one polymerizable dihydrogen phosphate-functional monomer is selected from the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate, and mixtures of the dihydrogen phosphate monoester of 2-hydroxyethyl methacrylate and the phosphoric acid diester of 2-hydroxyethyl methacrylate.
 - 37. A pigment slurry according to claim 35 wherein the at least one polymerizable dihydrogen phosphate-functional monomer is selected from the dihydrogen
- vinylbenzylphosphates, and CH₂=CCH₃CO₂CH₂CH₂OPO (OH)₂,

 CH₂=CCH₃CO₂CH₂CH [OPO (OH)₂] CH₃, CH₂=CCH₃CO₂CH₂CH₂CH₂OPO (OH)₂ and

 CH₂=CCH₃CO₂CH₂CHOHCH₂OPO (OH)₂
 - 38. An aqueous dispersion of composite particles, the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including polymerized residues of itaconic acid.
 - 39. An aqueous dispersion according to claim 38, the polymeric latex particles being polymerized from monomer including itaconic acid.
 - 40. A coating composition comprising an aqueous dispersion of composite particles, the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including polymerized residues of itaconic acid.
 - 41. A pigment slurry comprising an aqueous dispersion

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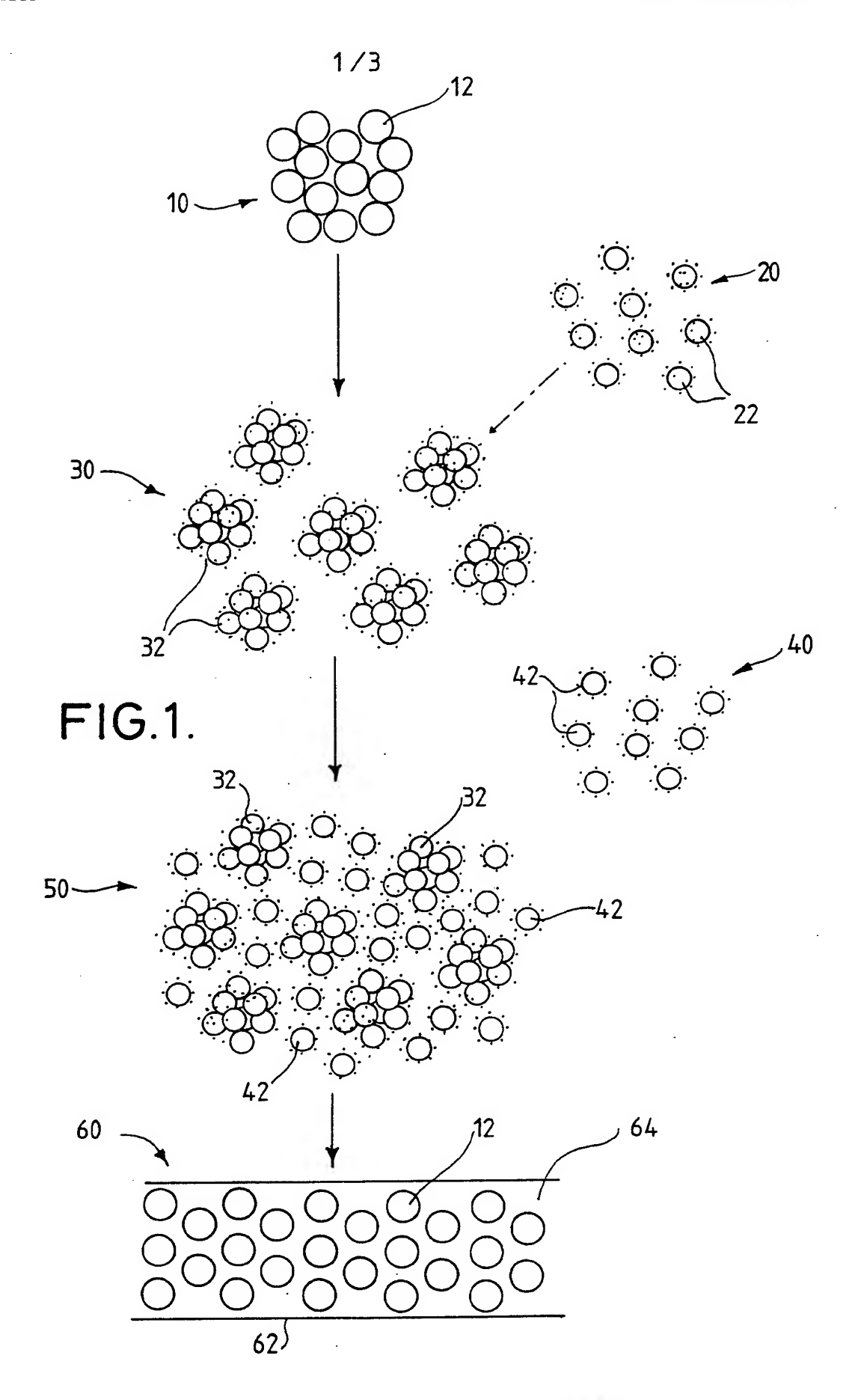
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of composite particles, the composite particles each comprising a plurality of polymeric latex particles adsorbed to a titanium dioxide particle, the polymeric latex particles including polymerized residues of itaconic acid.

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- 42. A process for preparing an aqueous dispersion of composite particles, the composite particles each comprising a plurality of selected polymeric latex particles adsorbed onto an inorganic material particle, the process comprising:
- a) dispersing inorganic material particles in an
 10 aqueous medium;
 - b) preparing a dispersion or suspension of selected polymeric latex particles in the aqueous medium, the polymeric latex particles being prepared by an emulsion polymerization process employing a monomer mixture including at least one polymerizable ethylenically unsaturated acid-functional monomer selected from the class consisting of monobasic and polybasic monomers having acidic functional groups including at least one proton with an acid equilibrium constant (pKa) of from about 4 to 8; and
 - c) mixing the aqueous medium containing the inorganic material particles and the selected polymeric latex particles, the selected polymeric latex particles adsorbing onto the inorganic material particles to provide the composite particles.
- 43. A process according to claim 42 wherein the acid equilibrium constant is about 6.



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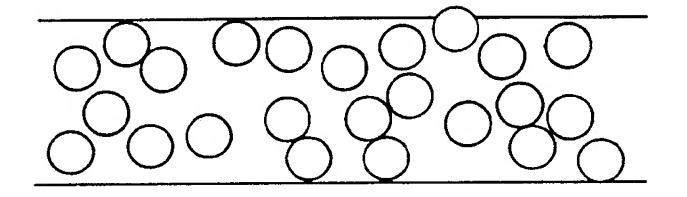


FIG. 2.

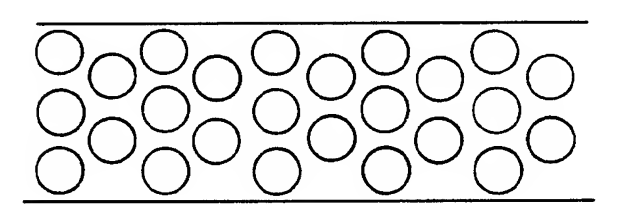


FIG. 3.

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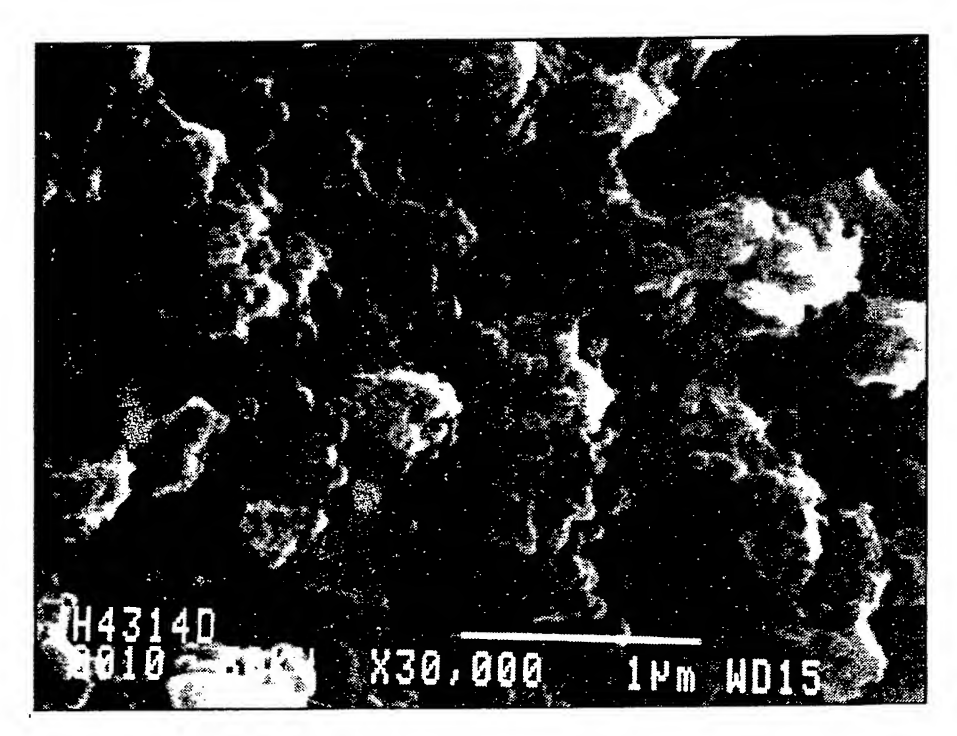


FIG.4a.

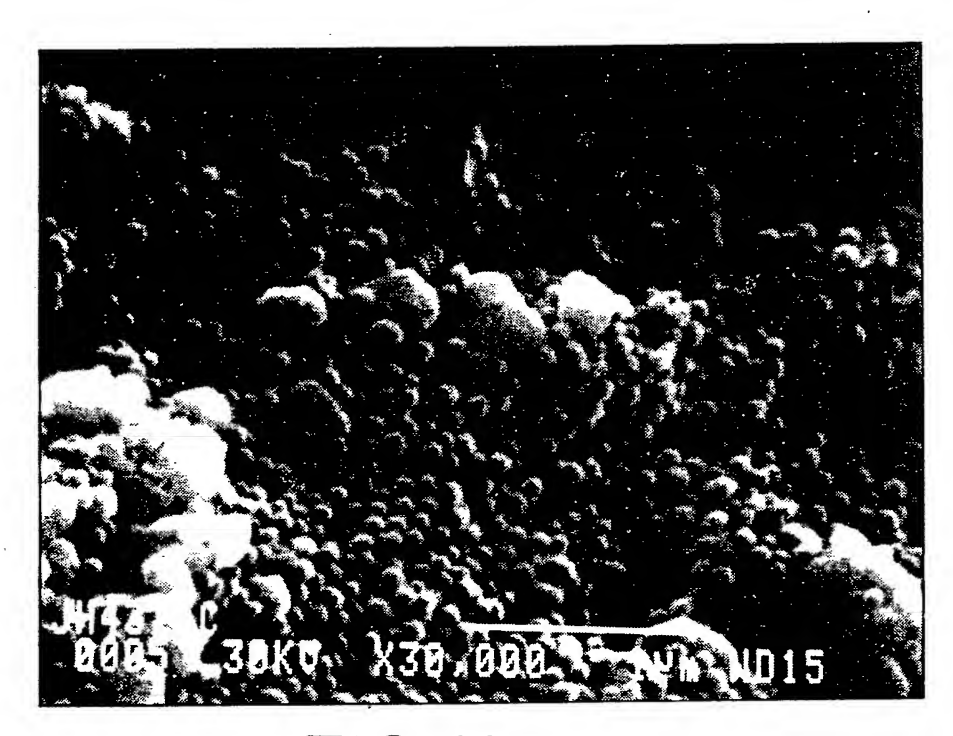


FIG.4b.

SUBSTITUTE SHEET

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 92/09648

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) ⁶							
According to Internation Int.C1. 5 CO9	nal Patent Classification (IPC) or to both Nati D5/02; C09D7/12;	onal Classification and I CO9C	PC 1/00;	C09D17/00			
II. FIELDS SEARCHED							
Minimum Documentation Searched ⁷							
Classification System Classification Symbols							
Int.Cl. 5	C09C ; C09D						
	Documentation Searched to the Extent that such Docu	other than Minimum Doments are Included in the	ocumentation Fields Searched®				
	NSIDERED TO BE RELEVANT ⁹		12	Relevant to Claim No.13			
Category Cita	ation of Document, 11 with indication, where a	propriate, of the relevant	passages	Relevant to Claim 140			
24	,A,4 025 483 (A. RAMIG, May 1977 e column 1, line 45 - line			1,2,4,5, 8-15,19, 20,41,42			
se 1- se							
	A,0 337 672 (CROWN DECOMOCTOBER 1989	RATIVE PRODUCT	TS)				
	,A,O 421 238 (HOECHST) April 1991	_					
•	,A,O 113 435 (DESOTO) July 1984						
"Special categories of cited documents: 10 "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed "V" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "A" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "A" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered novel or cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed inve							
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

9209648 US 67022 SA

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report.

The members are as contained in the European Patent Office EDP file on

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

05/0 05/04/93.

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US-A-4025483	24-05-77	None			
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EP-A-0421238	10-04-91	DE-A- CA-A- JP-A- US-A-	3932816 2026454 3124774 5135970	11-04-91 31-03-91 28-05-91 04-08-92	
EP-A-0113435	18-07-84	US-A- US-A- US-A- CA-A- JP-A-	4474909 4474910 4474911 1226400 59115367	02-10-84 02-10-84 02-10-84 01-09-87 03-07-84	

【公報種別】特許法第17条第1項及び特許法第17条の2の規定による補正の掲載 【部門区分】第3部門第3区分

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【出願番号】特願平5-510901

【国際特許分類第7版】

CO8F 2/22 MBS

[FI]

COSF 2/22 MBS

手 続 補 正 書

平成11年11月12日

特许广長官一股

1 事件の表示 平成5年特許購第510901号

2 発明の名称 重合体ラテックスを含有する複合体粒子の水性分散物

3 補正をする者

| 影件との開係 | 特許出額人

ローム アンド ハース カンパコー 氏名 (名称)

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電話 03 (3288) 1801

5 補正対象書類名 明細書及び請求の範囲翻訳文

6 補正対象項目名 明細書及び請求の範囲

7 補正の内容

- () 請求の範囲を別紙の通り補正する。
- (2)明細書第5-2 真第7 行の次に、以下を新行にて追加する。

「本発明はその実施閣様として以下を包含する。

- 1. 複合体粒子で、失々、複数の選択された重合体ラテックス粒子が一つの無 機材料粒子上に吸着したものからなる複合体粒子の水性分散物を製造する方法に
 - a 》 水性媒体中に無機材料粒子を分散し、
- b) 水性媒体中に選択された重合体ラテックス粒子の分散物又は懸濁物を 調製し、然も、前記重合体ラテックス粒子はエマルジョン重合法により製造され ており、該エマルジョン重合法は、
- 1) 少なくとも一種類の重合可能なエチレン系不飽和酸官能性単量体を 合む単量体混合物を用いたエマルジョン重合法であって、然も、前記少なくとも 一種類の重合可能なエチレン系不飽和酸官能低単量体が、

A)イタコン酸、及び

B)重合可能なオレフィン基を有するアルコールの燐酸二水紫エス テル、

からなる群から選択されているエマルジョン重合法;

- 2) 重合体ラテックス粒子の表面近傍に酸官能性を優先的に与えるエマ ルジョン重合法;及び
- 3) 少なくとも一種類のエチレン系不飽和酸官能性単量体を含む単量体 混合物を用いたエマルジョン重合法で、前記エチレン系不飽和酸官能佐単量体の 全重量が重合体ラテックス固体の少なくとも約10重量%を占めるエマルジョン 重合法;

からなる群から選択されており、

そして

c) 前記無機材料粒子を含有する水性媒体と前記選択された重合体ラテッ

クス粒子とを混合し、前記選択された重合体ラテックス粒子を前記無機材料粒子 上に吸着させて複合体粒子を与える、

- ことからなる、複合体粒子水性分散物製造方法。
- 2.無機材料粒子を顔料分散剂を用いることにより水性媒体中に分散し、前記 顔料分散剤が無機材料粒子上に吸着し、透択された重合体ラテックス粒子が前記 顔料分散剤の存在下で前記無機材料粒子上に吸着し、前記無機材料粒子を含む水 性媒体と前記選択された重合体ラテックス粒子とを混合した時、複合体粒子を与 える、前記1項に記載の方法。
- 3. 燐酸二水素エステルが、2ーヒドロキシエチルメタクリレートの燐酸モノエステル、2ーヒドロキシエチルメタクリレートの燐酸ジエステル、3ーヒドロキシプロビルメタクリレートの燐酸モノエステル、及び3ーヒドロキシプロビルメタクリレートの燐酸ジエステルから選択される。前計2項に記載の方法。
- 4. 顔科分散剤が高分子電解質である、前記2項に記載の方法。
- 5、願料分散剤が、ボリ(メタ)アクリル酸、ボリ(メタ)アクリル酸塩、(メタ)アクリル酸の高分子電解質共重合体、及び(メタ)アクリル酸の高分子電解質共重合体の塩から選択される前記 4 項に記載の方法。
 - 6、顔科分散剤が無機高分子電解質である、前記4項に記載の方法。
- 7. 顔料分散剤がポリ燐酸塩である、前記6項に記載の方法。
- 8. 選択された重合休ラテックスが、約20℃より高い有効ガラス転移温度を 有する、前記2項に記載の方法。
- 9. 選択された重合体ラテックスが、約50℃より高い有効ガラス転移温度を 有する、前記8項に記載の方法。
- 10. 重合体ラテックス粒子の平均粒径が、約20 n mから無機材料粒子の平均粒径の約4倍である、前記2項に記載の方法。
- 11. 重合体ラテックス粒子の平均粒径が、無機材料粒子の平均粒径の約半分から前記無機材料粒子の平均粒径にほぼ等しい、前記10項に記載の方法。
- 12.無機材料粒子の平均粒径が約100nm~10,000nmであり、重合体ラテックス粒子の平均粒径が約20nm~5000nmである、前記2項に

記載の方法。

- 13. 無機材料粒子の平均粒径が約200 nm~250 nmであり、重合作ラテックス粒子の平均粒径が約20 nm~1000 nmである、前記12項に記載の方法。
- 14. 選択された重合体ラテックスの室刻ガラス転移温度が約20℃より低く、前記選択された重合体ラテックスの平均積径が約80 n m ~ 600 n m である前記2項に記載の方法。
- 15. 選択された重合体ラテックスの平均粒径が約100 mm~400 mmである前記14項に記載の方法。
- 16. 選択された重合体ラテックス粒子の平均粒径が約30 nm~100 nm である前記8項に記載の方法。
- 17、選択された平均粒役が約50 n m ~ 80 n m である前記 16 項に記載の方法。
- 18. 無機材料が二酸化チタンである前記2項に記載の方法。
- 19. 前記2項に記載の方法により製造された被覆組成物。
- 20. 複合体粒子で、失々、複数の環状された重合体ラテックス分散剤粒子が一つの無機材料粒子上に吸着したものからなる分散された複合体粒子を含む水性分組成物を製造する方法において、
- a) 顕料分散剤を用いて永佳媒体中に無機材料粒子を分散し、その最初の 類料分散剤が前記無機材料粒子上に吸着され、
- b) 水性媒体中に選択された重合体ラテックス粒子を入れた分散物又は懸 衝物を調製し、然も、前記重合体ラテックス粒子はエマルジョン重合法により製 造されており、該エマルジョン重合法は、
- 1) 少なくとも一種類の重合可能なエチレン系不飽和酸官能性単量体を 含む単量体混合物を用いたエマルジョン重合法であって、然も、前記少なくとも 一種類の重合可能なエチレン系不飽和酸官能性単量体が、
 - A) イタコン酸、及び
 - B) 重合可能なオレフィン港を有するアルコールの燐酸二水素エス

ゔル、

からなる群から選択されているエマルジョン重合法:

- 2) 重合体ラテックス粒子の表面近傍に敲官龍性を優先的に与えるエマルジョン重合法: 及び
- 3) 少なくとも一種類のエチレン系不飽和酸官能性単量体を含む単量体 退合物を用いたエマルジョン重合法で、前記エチレン系不飽和酸官能性単量体の 全重量が重合体ラテックス固体の少なくとも約10重量%を占めるエマルジョン 重合法;

からなる群から選択されており、

そして

- c) 的記無機材料粒子を含有する水性媒体と前記選択された重合体ラテックス粒子とを混合し、前記選択された重合体ラテックス粒子を前記顔料分散剤の存在下に前記無機材料粒子上に吸着させて複合体粒子を与える、
- ことからなる、複合体粒子水低分散物製造方法。
- 21. 爆酸二水素エステルが、2ーヒドロキシエチルメタクリレートの燐酸モノエステル、2ーヒドロキシエチルメタクリレートの燐酸ジエステル、3ーヒドロキシプロピルメタクリレートの燐酸モノエステル、及び3ーヒドロキシプロピルメタクリレートの燐酸ジエステルから選択される、前記20項に記載の方法。
- 22. 重合体ラテックス結合剤粒子の水性分散物と、分散した複合体粒子を含む水性媒体とを混合することを更に含む、前記21項に記載の方法。
- 23. 複合体粒子で、夫々、撥酸二水素エステル官能基を有する複数の重合体 ラテックス粒子が二酸化チタン粒子に吸着されたものからなる複合体粒子の水性 分散物。
- 24. 重合体ラテックス粒子が、少なくとも一種類の重合可能な機能工水素官 能性単量体を含む単量体から重合されたものである。前記23項に記載の水性分 散物。
- 25.少なくとも一種類の重合可能な機酸二水素官能性単量体が、重合可能な ビニル基及び重合可能な非ビニルオレフィン基から選択された重合可能な基を有

するアルコールの燐酸三水素エステルから選択されたものである。前記 2 4 項に 記載の水性分散物。

- 26. 少なくとも一種類の重合可能な燐酸二水素盲能性単量体が、2ーモドロキシエチルメタクリレートの燐酸二水素モノエステル、及び2ーモドロキシエチルメタクリレートの燐酸二水素モノエステルと2ーモドロキシエチルメタクリレートの燐酸ジエステルとの混合物から選択されたものである。前記25項に記載の水性分散物。
- 28. 複合体粒子で、失々、複数の重合体ラテックス粒子が一つの二酸化チタン粒子に吸着したものからなる複合体粒子の水性分散物を含み、然も、前記重合体ラテックス粒子が燐酸二水素エステル宮能器を有する被覆期成物。
- 29. 重合体ラテックス粒子が、少なくとも一種類の重合可能な機能工水器官能性単量体を含む単量体から重合されたものである。前記28項に記載の被覆組成物。
- 30.少なくとも一種類の重合可能な燐酸二水素官能性単量体が、重合可能な ビニル基及び重合可能な非ビニルオレフィン基から選択された重合可能な基を有 するアルコールの燐酸二水素エステルから選択されたものである、前記30項に 記載の被覆組成物。
- 31. 少なくとも一種類の重合可能な機能二水素官能性単量体が、2--ヒドロキシエチルメタクリレートの機能二水素モノエステル、及び2--ヒドロキシエチルメタクリレートの機能二水器モノエステルと2--ヒドロキシエチルメタクリレートの機能ジエステルとの混合物から選択されたものである、前部29項に記載の被覆組成物。

- 33. 複合体粒子で、失々、複数の重合体ラテックス粒子が一つの二酸化チタン粒子に吸着されたものからなる複合体粒子の水性分散物を含み、燃も、前記重合体ラテックス粒子が構酸二水業エステル官能基を有する顔料スラリー。
- 34、重合体ラテックス粒子が、少なくとも一種類の重合可能な燐酸二水素官 能性単量体を含む単量体から重合されたものである、前記33項に記載の顔料ス ラリー。
- 35.少なくとも一種類の重合可能な燐酸二水素官能性単量体が、重合可能なビニル基及び重合可能な非ビニルオレフィン基から選択された重合可能な基を有するアルコールの燐酸二水素エステルから選択されたものである。前記34項に記載の顔料スラリー。
- 36.少なくとも一種類の重合可能な解散工水祭官能性単量体が、2-ヒドロキシエチルメタクリレートの構設工水器モノエステル、及び2-ヒドロキシエチルメタクリレートの構設工水器モノエステルと2-ヒドロキシエチルメタクリレートの構設ジエステルとの混合物から選択されたものである。前記35項に記載の顔料スラリー。
- 37. 少なくとも一種類の重合可能な燐酸二水素官能性単量体が、燐酸二水素 ビニルベンジル、及びCH₂ = CCH₃ CO₂ CH₃ CH₃ OPO (OH)₂、 CH₂ = CCH₃ CO₂ CH₃ CH₄ OPO (OH)₃] CH₃、CH₂ = CC H₃ CO₂ CH₃ CH₄ OPO (OH)₃、及びCH₃ = CCH₃ CO₃ CH₄ CHOHCH₂ OPO (OH)₃ から選択されたものである、前記35項 に記載の顔料スラリー。
- 38. 複合体粒子で、夹々複数の重合体ラデックス粒子が一つの二酸化チタン

(別候)

請求の範囲

- 1. 複合体粒子で、夫々、複数の選択された重合体ラテックス粒子が一つの無 機材料粒子上に吸着したものからなる複合体粒子の水性分数物を製造する方法に おいて、
 - a) 水性媒体中に無機材料粒子を分散し、
- b) 水性媒体中に選択された重合体ラテックス粒子の分散物义は懸濁物を 調製し、然も、前記重合体ラテックス粒子はエマルジョン重合法により製造され ており、該エマルジョン重合法は、
- 1) 少なくとも一種類の重合可能なエチレン系不飽和酸官能性単量体を 合む単量体混合物を用いたエマルジョン重合法であって、然も、前記少なくとも 一種類の重合可能なエチレン系不飽和酸官能性単量体が、
 - A) イタコン酸、及び
- B) 重合可能なオレフィン基を有するアルコールの爆酸二水素エステル、
- からなる群から選択されているエマルジョン重合法:
- 2) 重合体ラテックス粒子の表面近傍に酸官能性を優先的に与えるエマルジョン重合法;及び
- 3) 少なくとも一種類のエチレン系不能和酸官能性単量体を含む単量体 混合物を用いたエマルジョン重合法で、前記エチレン系不能和酸官能性単量体の 全重量が重合体ラテックス固体の少なくとも約10重量%を占めるエマルジョン 重合法;
- からなる群から選択されており、
- そして
- c) 前記無機材料位子を含有する水性媒体と前記選択された重合体ラテックス粒子とを混合し、前記選択された重合体ラテックス粒子を前記無機材料粒子上に吸着させて複合体粒子を与える、
- ことからなる、複合体位子水佐分散物製造方法。

- 粒子に吸着したものからなる複合体粒子の水性分散物であって、しかも前記重合体ラテックス粒子がイタコン酸の重合された残器を有する複合体粒子水性分散物。 39. 重合体ラテックス粒子が、イタコン酸を含む単量体から重合されたものである、前記38項に記載の水性分散物。
- 40. 複合体粒子で、央々、複数の重合体ラテックス粒子が一つの工酸化チタン粒子に吸着したものからなる複合体粒子の水性分散物であって、しかも前記重合体ラテックス粒子がイタコン酸の重合された残毒を有する複合体粒子水性分散物を含む被覆組成物。
- 41. 複合体粒子で、夫々、複数の重合体ラテックス粒子が一つの二酸化チタン粒子に吸著したものからなる複合体粒子の水性分散物であって、しかも前記重合体ラテックス粒子がイタコン酸の重合された複葉を有する複合体粒子水性分散物を含む顔料スラリー。
- 42. 復合体粒子で、夫々、複数の選択された集合体ラテックス粒子が一つの 無機材料粒子上に吸着されたものからなる複合体粒子の水性分組成物を製造する 方法において、
 - a) 水性媒体中に無機材料粒子分散し、
- b) 前記水性媒体中に選択された重合体ラテックス粒子を入れた分散物义は懸酒物を調製し、然も、前記重合体ラテックス粒子は、約4~8の酸平衡定数(pKa)を有する少なくとも一つのプロトンを含む酸性官能基を有する一塩基性及び多塩基性単量体からなる群から選択された少なくとも一種類の単合可能なエチレン系不飽和酸官能性単量体を含む単量体混合物を用いたエマルジョン重合法により調製されており、そして
- c) 前記無機材料粒子を含む水性媒体と前記選択された重合体ラテックス 粒子とを混合し、前記選択された重合体ラテックス粒子を前記無機材料粒子上に 吸着させて複合体粒子を与える、
- ことからなる、複合体粒子水性分散物の製造方法。
- 43、酸平衡定数が約6である前記42項に記載の方法。」

- 2. 無機材料粒子を顔料分散剤を用いることにより水性媒体中に分散し、前記 顔料分散剤が無機材料粒子上に吸着し、選択された重合体ラテックス粒子が前記 顔料分散剤の存在下で前記無機材料粒子上に吸着し、前記無機材料粒子を含む水 性媒体と前記選択された重合体ラテックス粒子とを混合した時、複合体粒子を与 える、前求項1に記載の方法。
 - 3. 顔料分散剤が高分子電解質である、請求項2に記載の方法。
- 4. 無機材料が二酸化チタンである請求項2に記載の方法。
- 5. 請求項2に記載の方法により製造された被覆組成物。
- 6. 複合体粒子で、夫々、複数の選択された重合体ラテックス分散剤粒子が一つの無機材料粒子上に吸着したものからなる分散された複合体粒子を含む水性分組成物を製造する方法において、
- a) 願得分散剤を用いて水性媒体中に無機材料粒子を分散し、その最初の 顔料分散剤が前記無機材料粒子上に吸着され、
- b) 水性媒体中に選択された重合体ラテックス粒子を入れた分散物又は懸 濁物を調製し、続も、前記重合体ラテックス粒子はエマルジョン重合法により製 造されており、該エマルジョン重合法は、
- 1) 少なくとも一種類の重合可能なエチレン系不飽和酸官能性単量体を 含む単量体混合物を用いたエマルジョン重合法であって、然も、前記少なくとも 一種類の重合可能なエチレン系不飽和酸官能性単量体が、
 - A) イタコン酸、及び
- B) 重合可能なオレフィン基を有するアルコールの機酸二水紫エステル、
- からなる群から選択されているエマルジョン重合法;
- 2) 重合体ラテックス粒子の表面近傍に酸官能性を優先的に与えるエマルジョン重合法;及び
- 3) 少なくとも一種類のエチレン系不飽和酸官能性単量体を含む単量体 混合物を用いたエマルジョン重合法で、前記エチレン系不飽和酸官能性単量体の 全重量が重合体ラテックス固体の少なくとも約10重量%を占めるエマルジョン

重合法:

からなる群から選択されており、

そして

- c) 前記無機材料粒子を含有する水性媒体と前記選択された重合体ラテックス粒子とを混合し、前記選択された重合体ラテックス粒子を前記顔科分散剤の存在下に前記無機材料粒子上に吸着させて投合体粒子を与える、ことからなる、複合体粒子水性分散物製造方法。
- 7. 複合体粒子で、夫々、燐酸二水紫エステル官能基を有する複数の重合体 ラテックス粒子が二酸化チタン粒子に吸着されたものからなる複合体粒子の水性 分散物。
- 8. 重合体ラテックス粒子が、少なくとも一種類の重合可能な燐酸二水素官 能性単量体を含む単量体から重合されたものである、請求項7 に記載の水性分 散物。
- 9. 少なくとも一種類の重合可能な燐酸二水素質能性単量体が、重合可能な ビニル基及び重合可能な非ビニルオレフィン基から選択された重合可能な基を有 するアルコールの燐酸二水素エステルから選択されたものである、請求項8 に 記載の水性分散物。
- 10. 複合体粒子で、夫々、複数の重合体ラテックス粒子が一つの二酸化チタン粒子に吸着したものからなる複合体粒子の水性分散物を含み、然も、胸記重合体ラテックス粒子が燐酸二水素エステル官能基を有する被覆組成物。
- 1.1、複合体粒子で、失々、複数の重合体ラテックス粒子が一つの二酸化チタン粒子に吸着されたものからなる複合体粒子の水性分散物を含み、燃も、前記重合体ラテックス粒子が燐酸二水素エステル官能基を有する顔料スラリー。
- 1.2. 複合体粒子で、失々複数の重合体ラテックス粒子が一つの二酸化チタン 粒子に吸着したものからなる複合体粒子の水性分散物であって、しかも前記重合 体ラテックス粒子がイタコン酸の重合された残塞を有する複合体粒子水性分散物
- 13. 複合体粒子で、夹々、複数の選択された重合体ラテックス粒子が一つの 無機材料粒子上に吸着されたものからなる複合体粒子の水性分組成物を製造する

方法において、

- a) 水性媒体中に無機材料粒子分散し、
- b) 前記水性媒体中に選択された重合体ラテックス粒子を入れた分散物又は懸濁物を調製し、然も、前記重合体ラテックス粒子は、約4~8の酸平衡定数(pKa)を有する少なくとも一つのプロトンを含む酸性官能基を有する一塩基性及び多塩基性単量体からなる群から選択された少なくとも一種類の重合可能なエチレン系不飽和酸官能性単量体を含む単量体混合物を用いたエマルジョン重合法により調製されており、そして
- c) 前記無機材料粒子を含む水性媒体と前記選択された重合体ラテックス 粒子とを混合し、前記選択された重合体ラテックス粒子を前記無機材料粒子上に 吸着させて複合体粒子を与える、
- ことからなる、複合体粒子水性分散物の製造方法。
- 14. 酸平衡定数が約6である請求項13に記載の方法。